#### **SECOND NOTICE OF DEFICIENCIES (NOD)**

HWMA/RCRA PART B PERMIT APPLICATION, VOLUME 14
FOR STORAGE AND TREATMENT UNITS AT THE
IDAHO NUCLEAR TECHNOLOGY AND ENGINEERING CENTER (INTEC)
LIQUID WASTE MANAGEMENT SYSTEM (ILMWS)
IDAHO NATIONAL ENGINEERING AND ENVIRONMENTAL LABORATORY
EPA ID No. ID4890008952

The following list of deficiencies was compiled by the Idaho Department of Environmental Quality (DEQ). The list identifies deficiencies found in Volume 14, Revision 1, Books 1 through 4 of the INEEL Part B Permit Application. While a specific form for a Permit Application does not exist, the list of deficiencies is organized in accordance with the RCRA Part B Checklist.

#### **GENERAL COMMENTS**

1. It appears that not all of the waste streams managed by the ILWMS have been identified in the response to NOD #1 or the first paragraph of Section C-1. If the list of "typical waste streams" is not complete, the revised Part B Permit Application must include the rest of the waste streams. Otherwise, the Department of Energy (DOE) must demonstrate that the list is representative of the wastes managed by the ILWMS.

#### **RESPONSE:**

The list of waste streams managed by the INTEC Liquid Waste Management System (ILWMS), provided in the response to the first Notice of Deficiency (NOD), represents all wastes currently treated in that system. However, other waste streams not currently identified may be processed by that system in the future, provided that they meet the ILWMS waste acceptance criteria and process tolerance limits identified in Sections C-2a(1) and D-8b(5) of this Part B Permit Application.

The first paragraph of Section C-1 of the Permit B Permit Application was revised to read as follows:

"The INTEC units described in this permit application are used to manage a variety of wastes generated from INEEL activities. Waste streams managed by the ILWMS include:

- Liquids generated incidental to conducting debris treatment, decontamination, and descaling activities on INEEL equipment, piping, and valves
- Rain water and snow melt that infiltrate into sumps and other containment areas
- Water from radioactive fuel storage basins and pools
- Mop water and other cleaning liquids generated incidental to cleanup activities conducted in radiological areas
- Analytical residues, excess samples, and expired analytical standards generated by sampling and analytical laboratory activities

- Solutions from preventative maintenance and corrective maintenance leak tests on process piping and valves
- Aqueous service wastes, such as steam condensate
- ILWMS treatment residuals that may require further processing
- Other waste streams not currently identified that conform to the ILWMS waste acceptance criteria and process tolerance limits identified in Sections C-2(a)(1) and D-8(b)(5), respectively."
- 2. DOE has not sufficiently addressed the requirements in IDAPA 58.01.05.008 and .012 [40 CFR §§ 264 Subpart AA and 270.24]. In addition to the process vents associated with the deep tanks, evaporators, and fractionators, the revised Part B Permit Application must include all sparged tanks (continuous or intermittent) equipped with process vents. The regulations in IDAPA 58.01.05.008 [40 CFR 264 Subpart AA] do not differentiate whether a process vent emits continuously or intermittently. The Part B Permit Application, Revision 1, has also failed to demonstrate whether the <u>cumulative</u> organic emissions from <u>all affected process vents</u> can be maintained below 3 lb/hr or 3.1 tons/yr. This limit cannot be exclusively utilized for a single vent. DOE must revise the total organic emission determination.

#### **RESPONSE:**

The only process vent associated with the units identified in the Part B Permit Application is the INTEC main stack. The tanks included in the ILWMS are not equipped with process vents. Instead, emissions from these units are vented to either the Vessel Offgas System and/or the Process Offgas System, which eventually lead to the INTEC main stack.

The only sources of volatile and semi-volatile organics managed by the ILWMS are small quantities of laboratory wastes that are discarded to the CPP-601 Deep Tanks. Engineering Design File (EDF)-2432 was prepared and included as Attachment 1 to the first NOD response, which was submitted to the DEQ in December 2002. This EDF reviewed the inventory of organics sent from CPP-602, CPP-630, and CPP-684 to the Deep Tanks and determined that although the organic concentration in the Deep Tanks may exceed 10 ppmw, making 40 CFR 264 Subpart AA applicable to this system, associated emissions of volatile organic compounds (VOC) are much less than 3 lb/hr and 3.1 tons/yr.

Other INTEC processes may generate small quantities of non-volatile organics that are also managed by the ILWMS. These non-volatile organics are primarily generated through decontamination and debris treatment activities in CPP-659. Decontamination activities elsewhere at the INTEC may result in the discharge of additional small quantities of non-volatile organics to the ILWMS.

A review of the INEEL Chemical Management System for the past five years indicates that the following quantities of non-volatile organics are routinely managed in the ILWMS on an annual basis. Material Safety Data Sheets for these products are included as Attachment 1 to this NOD response.

• Alkaline Rust Remover (> 60% sodium hydroxide, 10-20% triethanolamine, 1-10% sodium gluconate, 1-10% diethanolamine); 600 lbs/yr

- Oxalic Acid (99.6% oxalic acid, 0.4% inert salts); 58 lbs/yr
- Radiacwash (> 85% mineral water, 6 % octylphenol, 5.7 % tetrasodium ethylenediamine, 2.9% citric acid); < 4 gal/yr
- Small quantities of oil/grease from decontamination/debris treatment of equipment.

The only sources of volatile and semi-volatile organics to the ILWMS are the Analytical Laboratories, via the CPP-601 Deep Tanks. EDF-2432, previously submitted, conservatively assumes that organics volatilize immediately and completely upon introduction to the Deep Tanks and determined the organic emissions to be 0.035 ton/yr, approximately two orders of magnitude lower than the 3.1 tons/yr limit. By assuming complete volatilization of organics in the Deep Tanks, this value includes any potential emissions that could occur from the Process Equipment Waste Evaporator (PEWE) or Liquid Effluent Treatment and Disposal (LET&D) facilities.

Because the New Waste Calcining Facility (NWCF) Evaporator Tank System (ETS) is considered a segment of the ILWMS treatment train and will be added as a modification to the final permit, its contribution to the overall emission of organics must also be considered. Included as Attachment 2 to this NOD response is "NWCF Evaporator Tank System 2001 Offgas Emissions Inventory," INEEL/EXT-02-00198, February 2002. This report provides offgas and liquid stream characterization of the ETS while processing typical wastes from the Tank Farm Facility during May and June, 2001. The analytical results from the offgas sampling event indicate that the hourly total emissions rate for all volatile and semi-volatile organic emissions was less than 0.02 lbs/hr or less than 0.09 ton/yr. Thus, the combined contributions from all units that manage volatile and semi-volatile organics in the ILWMS = 0.035 ton/yr + 0.09 ton/yr = 0.125 ton/yr. This value is considerably less than the 3.1 tons/yr limit prescribed in 40 CFR Part 264 Subpart AA.

3. The Risk Assessment of Potential Hazardous Air Pollutant Emissions from the ILWMS (Attachment 2, Book 1 of 4, Part B Permit Application) does not satisfy the requirements of IDAPA 58.01.05.008 [40 CFR § 264.601]. This regulation requires that permits for miscellaneous units contain such terms and provisions as necessary to protect human health and the environment. Unlike other hazardous waste management units that have technology based performance criteria (e.g., incinerators or landfills), the permit conditions and performance standards for miscellaneous units are based on the risk assessment and engineering judgment. A combination of [40 CFR §§ 264.601 (c) and 264.601(c)(5)] reads as follows:

"Prevention of any releases that may have adverse effects on human health or the environment due to waste migration in the air considering the existing quality of the air, including other sources of contamination and their cumulative impact on the air."

Fugitive emissions from fixed sources and stationary emission sources, regardless of their status under HWMA, must be considered under this regulation.

The impact of sources and their cumulative impact are to be assessed on a site by site (or area by area) basis. At some facilities, it may be that adding the separate screening level risk assessments is the most efficient means to consider the cumulative risk. At other facilities, it may be necessary to conduct a complex assessment of cumulative risk, evaluating realistic and/or actual (e.g., background) emission scenarios.

Considering the complicated nature and variables associated with the INEEL, DEQ recommends that the DOE develop a risk assessment work plan to support evaluation of cumulative impacts from emission sources. At a minimum, the work plan should identify the following:

- 1. Sources of contamination to be included in the risk assessment;
- 2. The air dispersion model to be used;
- 3. Exposure models, both human receptor scenarios and ecological receptors, to be evaluated;
- 4. Emission estimates for each contaminant from each emission source:
- 5. Meteorological data to be included in the dispersion model;
- 6. Toxicity data for each chemical identified in the emissions estimate;
- 7. Proposed human health protectiveness criteria for both carcinogenic and non-carcinogenic risks; and,
- 8. Proposed ecological protectiveness criteria.

In order to avoid unnecessary protraction of the permitting process, DEQ is strongly suggesting that the work plan be approved prior to actual assessment of risk.

#### **RESPONSE:**

Per 40 CFR § 264.601, miscellaneous units must be located, designed, constructed, operated, maintained, and closed in a manner that will ensure protection of human health and the environment. Protection of human health and the environment includes prevention of releases due to migration of waste constituents in the groundwater, surface water, or air.

Protection of groundwater and surface water are ensured since Buildings CPP-604, CPP-641, CPP-649, CPP-659 Annex, CPP-1618, and CPP-601 are fully enclosed buildings equipped with secondary containment and leak detection devices to prevent the release of hazardous and mixed waste constituents. Sections F-4b and F-4c of the Part B Permit Application provide information regarding protection against runoff and contamination of water supplies. This response specifically addresses an evaluation of the impacts to the air from operation of the ILWMS.

Based on discussions with the DEQ on May 21, 2003, "Risk Assessment of Potential Hazardous Air Pollutant Emissions from the INTEC Liquid Waste Management System," submitted as Attachment 2 to the first NOD response for Volume 14 in December 2002, is considered to be adequate for addressing incremental human health impacts for the public from ILWMS releases. This assessment calculates individual contaminant risk and hazard quotients (HQ) using risk or HQ per unit release factors determined from the New Waste Calcining Facility (NWCF) Screening Level Risk Assessment (SLRA). This method is technically appropriate if: 1) the ILWMS has the same source release geometry (e.g., stack height, flow rate) as that modeled for the NWCF SLRA; 2) the meteorological data used in the NWCF SLRA modeling are appropriate for the ILWMS; and 3) the exposure scenarios evaluated in the NWCF SLRA are appropriate for the ILWMS. All three of these conditions were evaluated and determined to be consistent for the ILWMS. Therefore, the method of calculating ILWMS impacts from NWCF impacts on a

contaminant-specific basis is a technically appropriate method for risk assessment of the ILWMS emissions.

Ecological risk assessments use modeled soil concentrations as a starting point for calculating impacts to ecological receptors. Modeled soil concentrations for specific contaminants are directly proportional to the contaminant release rates for a given time of exposure. Based on this, and conditions 1-3 described above, ecological impacts for the ILWMS may be calculated in the same manner as human health effects, i.e., multiplying the ecological receptor HQ per g/s released from the NWCF Screening Level Ecological Risk Assessment (SLERA) by the ILWMS contaminant emission rates and then summing the HQs to obtain a total hazard index (HI).

Per 40 CFR § 264.601(c)(5), only sources that release pollutants to the air will be evaluated for "their cumulative impact on the air." The evaluation will exclude incremental impacts from non-air pathway sources (e.g., exposure via soil ingestion or absorption directly from buried solid waste). For the INEEL CERCLA Disposal Facility (ICDF), this would include the airborne emissions calculated in the Short-term Risk Assessment. It would not consider the impacts calculated for ecological receptors in the Screening Level Ecological Risk Assessment (SLERA) for the ICDF, because these impacts were calculated based on direct exposure to calculated design inventory landfill waste concentrations, without transport to the air. Further, the impacts calculated in the ICDF SLERA are not appropriate for use in this evaluation because the ICDF SLERA used very conservative (maximum design inventory) landfill waste concentrations and specifically states that the analysis was developed to support facility design only and should not be used to approximate actual site conditions.

A revised risk evaluation for ILWMS emissions is proposed, which will include the following:

- 1. A summary of the human health and ecological risk calculations from ILWMS emissions using the above-described methods.
- 2. Discussion on the relative importance of evaluating cumulative impacts on human health for <u>determination of ILWMS permitting.</u> The human impacts from ILWMS emissions (risk = 7E-08 and HI = 0.0024) are less than 1/100 (< 1%) of the currently used human health criteria for RCRA risk assessments (1E-05 risk or 0.25 HI). The potential ecological impacts are also likely a very small fraction of the accepted ecological criteria based on the results of the NWCF SLERA and the fact that the ILWMS emission rates are much smaller than the NWCF emission rates. Assessment of incremental impacts from existing sources becomes much less important when the impacts from the source evaluated are very small compared to acceptable impact criteria. When the impacts from the incremental source are 1% or less of the criteria, evaluation of existing source impacts has two likely outcomes: 1) the impacts from the existing sources will not increase the cumulative impacts beyond the criteria; or 2) the impacts from the incremental source will not appreciably change the existing air quality. For example, if the existing ambient air quality results in a risk of less than or equal to 9.8E-06 (slightly less than the risk criteria of 1E-05), adding in the ILWMS risk (7E-08) gives a cumulative risk of 9.9E-06; still less than the risk criteria. If the existing air quality results in a risk of 1E-05 or greater (greater than or equal to the risk criteria), adding in the ILWMS risk gives a cumulative risk of 1.007E-05 (or less). This is quantitatively the same value as the existing air quality risk within 2 significant figures, which is the maximum number normally justified given the uncertainty in the risk assessment process.
- 3. <u>Summary of ICDF cumulative impacts on human health.</u> One of the major sources of concern relating to cumulative impacts with the ILWMS is the ICDF, which is currently

being constructed just southwest of the INTEC fence line. The maximum human health impacts to the public calculated for this facility (ICDF Short-term Risk Assessment) using very conservative (maximum design inventory) assumptions were determined to be low enough ( $\leq$  3E-08 risk and  $\leq$  0.01 HI) that their cumulative impacts with the ILWMS emissions are well less than the risk criteria (7E-08 + 3E-08 = 1E-07). The cumulative HI from the ICDF and ILWMS would be  $\leq$  0.0124, far less than the 0.25 HI criteria. These cumulative impacts will be summarized. No additional human health risk assessment is planned for ICDF.

- 4. Area screening for cumulative ecological impacts. Cumulative impacts from ILWMS and existing sources may become an issue for on-site ecological impacts because of the much closer proximity of the receptors to the sources (right outside facility fence lines). To address this concern, screening modeling of existing sources will be performed to determine the INEEL facilities whose plumes significantly overlap with the ILWMS (INTEC main stack) plume. For this initial modeling, the following five areas will be evaluated: 1) INTEC main stack; 2) ICDF ground-level releases (from the landfills and evaporation ponds); 3) TRA; 4) CFA; and 5) RWMC. These facilities were selected based on their location within the INTEC main stack plume footprint determined in the NWCF SLRA. For this initial screening modeling only, TRA, CFA, and RWMC will be evaluated using a single 10-m high centrally located point source (this is reasonable given the lack of a single large source at these facilities and their relatively large distance from the INTEC maximum impact location, located approximately 1 km southwest of INTEC). The latest EPA version of the ISC3 model will be used for the modeling with 5-years of National Oceanic and Atmospheric Administration (NOAA) meteorological data from the Grid 3 tower, located 2-km north of INTEC. Unit release annual average air concentration will be modeled and plotted (as isopleths). If a facility contributes less than 10% of the total relative air concentration (from all sources) at the ILWMS maximum on-site impact location (located outside area fence lines), then that facility will be excluded from further analysis (its plume dispersion pattern does not significantly overlap that from the ILWMS). If a facility contributes more than 10% of the total relative air concentration at the ILWMS maximum impact location (located outside area fence lines), then refined modeling will be done for quantified sources at that facility using actual source characteristics (e.g., location, release height, flow rates).
- 5. Evaluation of cumulative ecological impacts from contributing sources. If a facility contributes 10% or more of total impact at the ILWMS maximum impact location using the above screening modeling, the following additional analyses will be performed to evaluate cumulative ecological impacts at the maximum impact location for the INTEC main stack (same as that modeled for NWCF):
  - Emissions from the ILWMS and contributing sources will be evaluated for pollutants with existing emissions rate data (either calculated or measured) and reliable toxicological data. Radionuclides will not be evaluated.
  - Cumulative deposition rates from contributing facilities will be calculated using the ISC3 model.
  - Cumulative soil concentrations will be calculated using approved EPA human health risk assessment guidance.
  - Ecological impacts will be assessed using HQs calculated by dividing the modeled soil concentrations by ecologically-based screening levels (EBSLs), as described in the

"Guidance Manual for Conducting Screening Level Ecological Risk Assessments at the INEL," INEL-95/0190, 1995. The HQs will then be summed across all pollutants to obtain an HI. This method was used in both the NWCF and ICDF SLERAS.

- The proposed ecological protectiveness criterion for initial screening of impacts is an HI of 1.0. An HI of 1.0 is considered appropriate because of the conservative nature of the EBSL approach and because this assessment takes into account cumulative impacts.
- A qualitative uncertainty analysis will be included which discusses the uncertainty associated with ecological modeling assumptions and parameters values used in the assessment.
- 6. For certain pollutants (e.g., mercury), existing measurements may be used in lieu of modeled concentrations for existing source impacts if the model predictions show unacceptable impacts.

Upon receipt of DEQ concurrence with the approach presented above, the INEEL will complete the risk evaluation proposed. Transmittal of the final certified report to the DEQ is anticipated within 270 calendar days after receipt of concurrence.

#### **SPECIFIC COMMENTS**

#### C. WASTE CHARACTERISTICS

4. DEQ does not concur with the response to NOD #24. The PEWE and LET&D are mixed waste treatment units. While DEQ does not regulate the radioactive component of the waste, DOE is requesting alternate handling and sampling of the waste due to radiation concerns. DEQ cannot evaluate the validity of these requests without information on the radiological component (e.g. isotopes, hazards associated with radioactive materials, etc.) of the waste. Thus, information on the radiological component of the waste must be included in the revised Section C of the Permit Application.

#### **RESPONSE:**

The following description was added after the second paragraph of Section C-1 of the Part B Permit Application:

"Radionuclides that contribute the majority of the activity for wastes managed in the ILWMS include Y-90, Sr-90, Cs-137, Ba-137m, Pu-238, Sm-151, Pu-241, Pm-147, Eu-155, Eu-154, Pu-239, Am-241, Co-60, Ni-63, Cs-134, Sb-125, H-3, Pu-240, Tc-99, Cd-113m, Te-125m, Pa-233, Np-237, Eu-152, Zr-93, Cm-244, Fe-55, Nb-93m, Nb-94, Ru-106, Rh-106, Cs-135, U-234, Ce-144, and Pr-144. Units that comprise the ILWMS are capable of handling high-level, transuranic, and low-level radioactive wastes. Activities of typical wastes range from <20 nCi/g to 50,000 nCi/g. The exposure rates associated with these process solutions routinely exceed 100 mrem/hr and can pose a potentially serious hazard to workers at the INEEL if appropriate protective measures such as time, distance, and shielding are not applied. As a result the INEEL is requesting the use of alternate handling and sampling techniques as proposed in this permit application."

Specific waste characterization information, including radioactive waste analyses, is maintained in the operating record.

# C-1. Chemical and Physical Analyses: IDAPA 58.01.05.008 and .012 [40 CFR §§ 264.13(a) and 270.14(b)(2)]

5. Page C-5, line 3 through 4, appears to indicate that there are test methods equivalent to the methods set forth in the Subpart C of 40 CFR 261 approved by the Director of the Idaho DEQ. Clarify if these methods have actually been approved by the Director and revise the Part B Permit Application to include a brief description of each of the equivalent methods.

#### **RESPONSE:**

The analyses noted in this Part B Permit Application do not require variances/deviations from the test methods identified in Tables C-1 and C-2. The application language referenced on page C-5 of the Part B Permit Application is included to assure the DEQ that when such variances are necessary, appropriate equivalent method approval by the Director of the DEQ will be sought.

6. Page C-5, line 7 through 9, states that "With few exceptions, units that comprise the ILWMS manage land disposal restricted waste liquids that exhibit the characteristics of corrosivity and toxicity, and contain one or more listed constituents." In the revised Part B Permit Application, clarify the exceptions referred to in the sentence.

#### **RESPONSE:**

Information from lines 18 through 20 on page C-5 of the Permit B Permit Application was incorporated into the referenced paragraph to provide clarification. The description on page C-5 now reads:

"Except for the CPP-641 Westside Waste Holdup Tanks (VES-WL-103, VES-WL-104, and VES-WL-105), units that comprise the ILWMS manage land disposal restricted waste liquids that exhibit the characteristics of corrosivity and toxicity, and contain one or more listed constituents. Transfer lines from the Westside Waste Holdup Tanks include sections of tile-encased lines. Because of compatibility concerns regarding waste acids and the grout used for the tile-encasement, these tanks are prohibited from managing wastes exhibiting the characteristic of corrosivity (EPA HWN D002)."

7. The Part B Permit Application must provide detailed physical and chemical characteristics, based on analytical data and/or acceptable process knowledge, of waste being stored and/or treated in each of the ILWMS tank systems and miscellaneous units. For example, the Engineering Design File (EDF) for VOC Emissions from ALD Inputs to the INTEC Deep Tanks (see Attachment 1 of the Part B Permit Application, Revision 1), page 2 of 6, 3<sup>rd</sup> and 4<sup>th</sup> paragraphs, provide sufficient information pertaining to chemical characteristics of the waste being stored and treated in the CPP-601 Deep Tanks. The revised Part B Permit Application must provide equivalent quality of information/description for each unit associated with the ILWMS.

#### **RESPONSE:**

Attachments 3a through 3k to this NOD response contain analytical results for representative samples from tanks included in the ILWMS. Due to the level of redundancy designed into the system, several tanks and miscellaneous treatment units may manage the same waste. The following matrix is provided to show which analytical results are representative of wastes in the various segments of the ILWMS.

Tank(s)/Treatment Unit(s)	Description	Typical Analytical Results
VES-NCC-119	Fluoride Hot Sump Tank	See Attachment 3a
VES-NCC-122	Non-Fluoride Hot Sump Tank	See Attachment 3b
VES-NCD-123/VES-NCD-129	Decon Holdup and Collection Tanks	See Attachment 3c
VES-WL-132	CPP-604 Evaporator Feed Sediment Tank – fed through VES-WL-133	See Attachment 3d
VES-WL-133	CPP-604 Evaporator Feed Collection Tank	See Attachment 3d
VES-WL-102	CPP-604 Surge Tank for VES-WL-133 – fed through VES-WL-133	See Attachments 3d and 3e
VES-WL-109	CPP-604 Evaporator Head Tank – fed from VES-WL-133	See Attachment 3d
EVAP-WL-129 / EVAP-WL-161	Process Equipment Waste Evaporators – fed from VES-WL-133	See Attachment 3d
VES-WL-134	CPP-604 Process Condensate Surge Tank – may be used for series operation of the PEW evaporators or for storage capacity of concentrated acidic LET&D bottoms	See Attachment 3j or Attachment 3k
VES-WL-131	CPP-604 Process Condensate Surge Tank – feeds to the Process Condensate Collection Tanks	See Attachment 3j
VES-WL-108	CPP-604 Process Offgas Knock Out Pot – collected liquid would drain to either VES-WL-133 or VES-WL-131	See Attachment 3d or Attachment 3j

Tank(s)/Treatment Unit(s)	Description	Typical Analytical Results
VES-WL-101 VES-WL-111	CPP-604 Bottoms Collection Tanks – sample drawn from sample station WL-613 in 1983/Bottoms historically transferred to Tank Farm Tanks VES-WM-186	See Attachment 3f
VES-WL-103, VES-WL-104, and VES-WL-105	CPP-641 Westside Waste Holdup Tanks - these tanks have been emptied to the maximum extent allowed by the transfer pumps, without causing damage to the pump bearings, and no transfers of waste to this system are taking place	See Attachment 3g
VES-WM-100, VES-WM-101, and VES-WM-102	CPP-604 Tank Farm Tanks	See Attachment 3h
VES-WG-100, VES-WG-101, VES-WH-100, and VES-WH-101	CPP-601 Deep Tanks	See Attachment 3i
VES-WL-135, VES-WL-136, VES-WL-137, VES-WL-138, VES-WL-139, VES-WL-142, VES-WL-144, and VES-WL-150	Process Waste Liquid Tanks – may collect PEW evaporator condensate	See Attachment 3j
VES-WL-106, VES-WL-107, and VES-WL-163	CPP-604 Process Condensate Collection Tanks	See Attachment 3j
VES-WLK-197	CPP-1618 Acid Fractionator Waste Feed Head Tank – fed from CPP-604 Process Condensate Collection Tanks	See Attachment 3j
FRAC-WLL-170 and FRAC-WLK-171	CPP-1618 Acid Fractionators – fed from CPP-604 Process Condensate Collection Tanks	See Attachment 3j
VES-WLL-195	CPP-1618 Acid Fractionator Bottoms Tank	See Attachment 3k

Tank(s)/Treatment Unit(s)	Description	Typical Analytical Results
VES-NCR-171	CPP-659 Annex LET&D Nitric Acid Recycle Tank – fed from VES-WLL-195	See Attachment 3k
VES-NCR-173	CPP-659 Annex LET&D Nitric Acid Recycle Head Tank – fed from VES-NCR-171	See Attachment 3k

8. The response to NOD #29 states that "Sampling and analysis has demonstrated that when these small quantities of ignitable waste are aggregated with other waste in the CPP-601 Deep Tanks to facilitate treatment, the characteristic of ignitability is lost." In the absence of a summary of past studies, which have been conducted on wastes in the Deep Tanks, DEQ is unable to reach the same conclusion. The revised Section C must clearly justify the absence of the characteristic of ignitability in the deep tank.

#### **RESPONSE:**

As indicated in Table C-3 of the Part B Permit Application, process samples are taken from the CPP-601 Deep Tanks (VES-WG-100, VES-WG-101, VES-WH-100, and VES-WH-101) prior to each transfer to the PEWE system. These process samples are analyzed for flashpoint to ensure they do not exhibit the characteristic of ignitability. Attachment 4 to this NOD response contains templates for the parameters analyzed for process samples from the WG and WH tanks. This attachment also includes examples of results from recent process sampling activities from these tanks demonstrating that these mixtures are not ignitable.

9. Clarify whether or not the contents in the WWH tanks are corrosive.

#### **RESPONSE:**

Before the tanks were emptied to their current levels demineralized water was added until the pH of the waste was greater than or equal to 2.0 and less than or equal to 12.5 to ensure transferred waste was compatible with the grout used for tile-encased transfer lines. Attachment 5 to this NOD response shows analytical results of the material contained in the WWH tanks. These results show that the wastes in all 3 tanks exhibit a pH  $\geq$  2.0 and  $\leq$  12.5.

10. Results of the Balance of Plant sampling conducted in FY 1999 and 2000 (Book 1 of 4 of the Part B Permit Application) identify "unknown" volatile and semi-volatile organics (VOCs and SVOCs). The revised Section C must comprehensively discuss these unknown VOCs and SVOCs, include justifications as to why they were not/could not be identified, and discuss the fate of these unknown components throughout the ILMWS.

#### **RESPONSE:**

EPA guidance (Contract Laboratory Program National Functional Guidelines for Organic Data Review, EPA/R-94/012, 1994; Guidance on Collection of Emissions Data to Support Site-Specific Risk Assessments at Hazardous Waste Combustion Facilities, EPA530-D-98-002, August 1998; and USEPA Contract Laboratory Program Statement of Work for Organic Analysis, Multimedia, Multi-Concentration, OLM04.2, May 1999) specifies the identification and quantification of tentatively identified compounds (TICs) that are observed in the VOC and SVOC chromatograms. Results of the Balance of Plant sampling conducted in fiscal years (FY) 1999 and 2000 were reviewed by Analytical Laboratory personnel to identify TICs based on this guidance. Identification and quantification of these TICs was done according to the EPA Contract Laboratory Program guidelines. These guidelines indicate that chromatogram peaks with the greatest apparent concentrations should be tentatively identified and quantified. Additionally, all peaks with favorably matched retention times and apparent primary ion concentration down to 10% relative intensity (based on area) with respect to the primary ion of the nearest internal standard, should be identified and reported in their estimated concentration. Relative major ion intensities should agree within +/- 20%. Compound identifications were made using National Institute of Standards and Technology /EPA/National Institute of Health (May 1992) equivalent mass spectral library data. These are tentative identifications because there were no reference standards analyzed at the same chromatogram retention time as the tentatively identified compounds.

The reportable concentrations for these TICs were estimated by comparing the compound total area count (or peak height) to the total area count (or peak height) of the nearest internal standard free from interferences on the total or reconstructed ion chromatogram, and assuming a relative response factor (RRF) of 1.0. Results were reported on a separately identified list and flagged as estimated. The RRF is compound-specific, and cannot be determined in the TIC evaluation. The revised report specifies the corresponding internal standards used in the calculation of TIC concentrations.

For this effort, the "match quality" (Q), an agreement between the unknown peak and potentially matching library compounds, was required to be at least 85% for the unknown peak to be identified as a specific compound. This degree of matching may result in false-positive identification of TICs in the sample. Therefore, it may be necessary to re-examine results that, in the judgment of the project technical lead and quality assurance office, seem incredible. This list, however, will help identify any constituents that should be added to the VOC and SVOC target analyte list for future sampling. Any compounds that failed to meet a minimum match quality of 85% continue to be listed as "unidentified."

Based on the review of Balance of Plant analytical results for sampling conducted in FY 1999 and 2000, all reported compounds, with two exceptions, continue to be "unknown" (657 reported), "unknown hydrocarbons" (30 reported) or substituted benzenes (17 reported) in the referenced data packages since the relative major ion intensities exceeded the +/-20% criteria. The exceptions are as follows:

- Benzoic acid in VES-NCD-123 at an estimated level of 21 ug/L
- Chloroform in VES-WL-106 at an estimated level of 12 ug/L.

Both of these TICs are short-chain hydrocarbons that may be present in extremely low concentrations. EDF-2432 was prepared and included as Attachment 1 to the first NOD response, which was submitted to the DEQ in December 2002. This EDF reviewed the inventory of organics sent from CPP-602, CPP-630, and CPP-684 to the Deep Tanks. The EDF conservatively assumed that all "unidentified" volatile organic compounds were isopropanol, which was then readily converted to acetone; considered a worst-case volatile organic. The EDF concluded that although the organic concentration in the Deep Tanks may exceed 10 ppmw, making 40 CFR 264 Subpart AA applicable to this system, associated emissions of volatile organic compounds (VOC) are much less than 3 lb/hr and 3.1 tons/yr. The TICs and estimated concentrations identified from this review of FY 1999 and 2000 Balance of Plant sampling data do not alter this conclusion.

Provided as Attachment 6 to this NOD response are the analytical results identifying the TICs described above.

### C-lc. Waste in Miscellaneous Treatment Units: IDAPA 58.01.05.008 [40 CFR § 264.601(a)(1)]

11. Response to NOD #35 does not fully address DEQ's concern. The revised Section C must provide detailed physical and chemical characteristics, based on analytical data and/or acceptable process knowledge, of the evaporator and fractionator bottoms and overheads.

#### **RESPONSE:**

See the response to item No. 7 of this NOD.

# C-2. Waste Analysis Plan: IDAPA 58.01.05.008 and .012 [40 CFR §§ 264.13(b) and (c), and 270.14(b)(3)]

12. One of the objectives of the WAP (6<sup>th</sup> bullet) is to provide additional requirements for the characterization and acceptance of ignitable and reactive wastes. It is DEQ's understanding that the ILWMS will not manage wastes that exhibit the characteristic of reactivity (EPA HWN D003). Clarify this discrepancy in the revised Section C.

#### **RESPONSE:**

The DEQ is correct. The INEEL will not manage wastes exhibiting the characteristic of reactivity in the ILWMS. The 6<sup>th</sup> bullet under Section C-2 of the Part B Permit Application has been revised to read:

- "Provide additional requirements for the characterization and acceptance of ignitable wastes."
- 13. One of the examples of process knowledge described in Section C uses analytical reports from non-SW-846 chemical analyses, outdated chemical analyses, or information from similar processes. Justify the validity of these analyses being used as process knowledge in lieu of acceptable knowledge based on valid analytical techniques (EPA Guidance Manual for Waste Analysis at Facilities that Generate, Treat, Store and Dispose of Hazardous Wastes).

#### **RESPONSE:**

Non-SW-846 chemical analyses are process sampling results that do not necessarily follow SW-846 sample collection or QA/QC protocol prescriptively. The results, however, supply information related to the characteristics of the waste that may be used as process knowledge during RCRA characterization. If process sample results are inconsistent with the waste characterization information provided by the generator or indicate that the waste generating process may have changed, then the waste is recharacterized.

Outdated chemical analyses are not used to support process knowledge for RCRA characterization of wastes managed by the ILWMS. The reference to outdated chemical analyses has been removed from the Part B Permit Application.

The sixth bullet of Section C-2a of the Part B Permit Application was revised to read:

• "Analytical reports from non-SW-846 chemical analyses or information from similar processes."

#### C-2a(1). Waste Acceptance Criteria

14. State the concentration limits for both total suspended solids and total dissolved solids acceptable to the ILWMS.

#### **RESPONSE:**

There are no concentration limits for TSS or TDS. All physical and chemical operational constraints and tolerance limits are identified in Section D-8b(5) of the Part B Permit Application.

#### C-2c(1). Standard Sampling Methods

15. Samples from the ILWMS are typically collected through double hypodermic needle (double-needle) samplers, sample nozzles, or spigots. The double-needle samplers may lose VOCs and SVOCs to either headspace of the sample vial or to the carrier gases used to move/collect the sample. DOE must demonstrate that the use of the double-needle sampling system does not impact the accuracy of the VOC and SVOC analytical data.

### **RESPONSE:**

Appendix C-2 has been added to the Part B Permit Application. This appendix contains a report from Science Applications International Corporation entitled; "Final Report for Organics Partitioning Resulting from Operation of an INTEC Double-Needle Sampler, Revision 1," dated September 24, 2002. This report shows that volatile organics taken from a double-needle sampler closely correlate to samples taken from a spigot at a PEWE mock-up facility. In fact, in several instances, the level of volatile organics measured from samples taken through the double-needle sampler was slightly greater, or more conservative, than that for samples taken from the spigot.

The following description was added to the end of the first paragraph of Section C-2c(1) of the Part B Permit Application:

"Appendix C-2 contains a report from Science Applications International Corporation entitled, "Final Report for Organics Partitioning Resulting from Operation of an INTEC

Double-Needle Sampler, Revision 1," dated September 24, 2002. This study compares organic concentrations obtained from double-needle and spigot sampling techniques to determine whether potential stripping of organics occurs. The results of these tests indicate that INTEC sample collection and handling procedures do not significantly affect the concentration of volatile or semi-volatile organic constituents in the waste stream."

#### C-2c(3). Process Sampling

16. Exhibit C-1 of the Volume 14 Permit Application, Revision 0, shows two additional process sampling locations, the feed collection tanks and the bottom collection tanks, which are not identified in Table C-3 of the Permit Application, Revision 1. Clarify this discrepancy in the revised Section C.

#### **RESPONSE:**

Table C-3 in the Part B Permit Application identifies ILWMS typical process sampling locations and the parameters tested. Exhibit D-1, PEWE System Flow Diagram and Inputs, of the permit application has been revised to indicate that the Bottoms Collection Tanks and the PEWE Feed Tanks are not routinely sampled.

Bottoms collected in VES-WL-101 and VES-WL-111 are transferred to either the Tank Farm Facility or the Evaporator Tank System where sampling can occur. Since these tank systems were designed and constructed to manage the types of waste to be processed, there is no need to collect process samples prior to waste transfers.

VES-WL-102 and VES-WL-133 contain samplers; however, the sample lines are plugged and not serviceable. Process samples are taken upstream of the PEWE Feed Tanks to ensure that process tolerance limits are met. VES-WL-132 is not equipped with a sampler.

17. Justify why the analytical parameters seem to vary with the sampling location. Each waste stream must meet the same acceptance profile before it can be received to the evaporators.

#### **RESPONSE:**

Table C-3 identifies typical ILWMS process sampling locations and parameters. Process sampling differs from RCRA characterization sampling in that process samples are collected solely to ensure optimum operation of the miscellaneous treatment units. RCRA characterization and waste verification/acceptance are completed prior to receiving waste streams into the ILWMS. Process samples are taken after acceptance of the waste into the system to ensure performance criteria associated with the miscellaneous treatment units are satisfied for each consecutive step in the treatment train. Process samples are not taken for RCRA characterization of wastes. However, if process sampling results are inconsistent with the waste characterization information provided by the generator or indicate that the waste generating process may have changed, then the waste is recharacterized.

The following statement was added to the last paragraph of Section C-2c(3) of the Part B Permit Application.

"If process sampling results are inconsistent with the waste characterization information provided by the generator or indicate that the waste generating process may have changed, then the waste is recharacterized."

# C-2f. Additional Requirements for Ignitable, Reactive, or Incompatible Wastes: IDAPA 58.01.05.008 [40 CFR §§ 264.13(b)(6) and 264.171

18. It appears that a hexone/nitric acid reaction can occur if an adequate concentration of hexone is present and necessary temperature requirements are met. The flashpoint of hexone in water at the elevation of the INEEL is 133°F (56°C) at a concentration of 2000 mg/L. Since the operating temperature of the PEW evaporators and LET&D fractionators are high enough to sustain a hexone/nitric acid reaction, under the operating condition, the concentration of total organic carbon (TOC) allowed in the feed of the ILWMS is limited to 1100 mg/L (assuming all TOC is hexone). These descriptions are included in the NOD response (response #51) but not in the Part B Permit Application, Revision 1. Because the tolerance limit set for the TOC appears to be a requirement for reactive waste, include the response to NOD #51 to the revised Section C.

#### **RESPONSE:**

The following description was added to the end of Section C-2f of the Part B Permit Application:

"The safety analysis documentation for the ILWMS indicates that, under the proper conditions, two potentially explosive reactions could occur. These reactions are tributyl phosphate (TBP) with nitric acid and hexone with nitric acid. Due to the temperature requirements necessary for these reactions, the only units described in this Part B permit application that could potentially sustain these reactions are the PEW evaporators and the LET&D fractionators.

"Conditions necessary for a TBP/nitric acid reaction include appropriate TBP concentration and elevated temperature (studies have shown that this reaction does not become extremely exothermic until the solution reaches 186° C).

"The quantity of TBP in the ILWMS is extremely small. Since the end of fuel reprocessing activities at the INTEC in the early 1990's, no TBP has been added to the system. In addition, all liquids in the INTEC TFF have already been evaporated at least once, further reducing the volume of TBP.

"Both the PEWE and LET&D facilities operate at much lower temperatures than are necessary to sustain a TBP/nitric acid reaction. Standard operating temperatures for these facilities are 108° C for the PEW evaporators and 118° C for the LET&D fractionators. Deviations from these operating temperatures result in waste feed cutoffs and/or shutdown of the system well before a temperature of 186° C can be reached.

"In order for a hexone/nitric acid reaction to occur, similar conditions must exist. A reaction can only be sustained if an adequate concentration of hexone is present and necessary temperature requirements are met. The flashpoint of hexone in water at the elevation of the INEEL is 133° F (56° C) at a concentration of 2000 mg/L.

"Like TBP, hexone is present in INTEC liquid wastes in only minimal concentrations. However, since the operating temperatures of the PEW evaporators and LET&D fractionators are high enough to sustain a hexone/nitric acid reaction, under the

appropriate conditions, the concentration of TOC allowed in the feed to the ILWMS is limited to 1100 mg/L. To ensure conservatism, all TOC is assumed to be hexone. This tolerance limit is identified in Section D-8b(5) of this permit application for both the PEWE and LET&D facilities.

"Furthermore, the LET&D facility is operated as an open system. The LET&D fractionators are maintained at a pressure of –20 in. water column. Both the TBP/nitric acid and the hexone/nitric acid scenarios require a closed system to sustain a reaction. The conditions in the LET&D facility preclude either reaction from occurring. If a vacuum cannot be maintained, the treatment process is automatically shut down.

"Therefore, the risk of explosive TBP/nitric acid reactions is eliminated due to low reactant concentrations, inadequate temperature, and open vessel conditions in the LET&D fractionators. Similarly, hexone/nitric acid reactions cannot occur due to low reactant concentrations, which are further regulated by administrative controls, and open vessel conditions on the LET&D fractionators."

19. The last sentence of the response to NOD #53 should be included in the revised waste analysis plan.

#### **RESPONSE:**

The following description was added to the first paragraph of Section C-2f of the Part B Permit Application:

"As identified in Table C-3 of Section C-2c(3), process samples are taken from the CPP-601 Deep Tanks prior to each transfer to the PEWE system and tested for flashpoint to ensure the feed stream is not ignitable."

# C-4. Subpart AA, Subpart BB, and Subpart CC Applicability: IDAPA 58.01.05.008 [40 CFR §§ 264.1030, 264.1050, and 264.1080]

20. DEQ's comments appear to have not been adequately addressed and included in the Permit Application, Revision 1. See the General Comment section, second bullet.

#### **RESPONSE:**

See the response to item No. 2 of this NOD.

#### D. PROCESS INFORMATION

21. The response to NOD #67 indicates that PEWE off-gas samples, based on a study conducted on a bench-scale model of the PEWE, taken downstream of the PEWE condensers did not contain free liquids, thus the off-gas is non-condensable. The mist eliminator and superheater downstream of the PEWE condenser appear to be redundant equipment based on the bench-scale study. The revised Part B Permit Application must describe the need and operational parameters for this equipment.

#### **RESPONSE:**

The mist eliminator and superheater, components of the Vessel Offgas (VOG) System, are included to protect and extend the life expectancy of high-efficiency particulate air (HEPA) filters included in the offgas system. Under normal operations these components might be considered redundant; however, in the unlikely event of system upset they ensure adequate protection of human health and the environment by preventing degradation of the HEPA filters.

The first paragraph of Section D-2d of the Part B Permit Application under the heading, "PEWE Pressure Controls" was revised to read:

"Waste treatment and storage vessels at INTEC are connected to a gaseous waste treatment system called the VOG system. All ILWMS storage and treatment systems discharge gases such as instrument air purges (used in level, density, and pressure instrumentation), air spargers (agitators), and gases displaced from a vessel when it fills with liquid.

"Gases from the PEWE and other INTEC processes, such as the Tank Farm and CPP-659 vessels, vent to the CPP-604 VOG system. The CPP-604 VOG system consists of a mist eliminator, superheater, and HEPA filter banks. The mist eliminator and superheater are included to protect and extend the life expectancy of the HEPA filters.

"Several facility process and vessel offgas systems, including the CPP-604 VOG system, combine in the Process APS located in CPP-649. The Process APS is a back-up system that treats the combined process and vessel offgas streams from CPP-601, CPP-604, Tank Farm, and the NWCF. The Process APS treatment consists of a mist eliminator, superheater, and HEPA filters. Next, the process and vessel offgases are routed to the INTEC Main Stack (CPP-708) where they mix with building ventilation air and are exhausted to the atmosphere. The vessel and process offgas systems are maintained under a vacuum to control contamination. The system equipment and piping are fabricated from acid resistant stainless steel for corrosion resistance. Additionally, the Process Condensate Collection Tanks can be vented to the process condensate collection cells, which vent to the CPP-604 building ventilation system. The VOG and APS systems are described further in Section D-8b of this application."

22. To qualify for the emergency structure exemption as set forth in IDAPA 58.01.05.008 [40 CFR § 264.1(g)(8)(i)], the revised Section C must clarify the source(s) of the wastes collected in the PWL sumps.

#### **RESPONSE:**

Sump SU-WL-140 is located in the South Cell of the Rare Gas Plant (RGP). The RGP is no longer active. Therefore, there are no sources of waste that would be collected in this sump.

Sump SU-WL-143 is located in the RGP Pump Pit. Since the RGP is no longer active, there are no sources of waste that would be collected in this sump.

Sump SU-WL-148 is located at the INTEC main stack. In the event of equipment failure, condensate from the main stack could collect in this sump.

Sumps SU-WL-145 and SU-WL-146 are part of the secondary containment and leak detection system in the PEWE Condensate Collection Cell. As such, these sumps do not require the emergency structure exemption set forth in IDAPA 58.01.05.008 [40 CFR § 264.1(g)(8)(i)].

Sump SU-WL-147 is part of the secondary containment and leak detection system in the PEWE EVAP-WL-161 Cell. As such, this sump does not require the emergency structure exemption set forth in IDAPA 58.01.05.008 [40 CFR § 264.1(g)(8)(i)].

The following description was added to Section C-1b of the Part B Permit Application under the heading, "Process Waste Liquid (PWL) System (VES-WL-135, VES-WL-136, VES-WL-137, VES-WL-138, VES-WL-139, VES-WL-142, VES-WL-144, and VES-WL-150)":

"Sumps SU-WL-140, -143, -145, -146, -147 and -148 do not contain tanks. These sumps are not used routinely. The exclusive purpose of these sumps is to contain liquids during immediate responses to discharges of hazardous wastes.

"Sump SU-WL-140 is located in the South Cell of the Rare Gas Plant (RGP). The RGP is no longer active. Therefore, there are no sources of waste that would be collected in this sump.

"Sump SU-WL-143 is located in the RGP Pump Pit. Since the RGP is no longer active, there are no sources of waste that would be collected in this sump.

"Sump SU-WL-148 is located at the INTEC main stack. In the event of equipment failure, condensate from the main stack could collect in this sump.

"Sumps SU-WL-145 and SU-WL-146 are part of the secondary containment and leak detection system in the PEWE Condensate Collection Cell.

"Sump SU-WL-147 is part of the secondary containment and leak detection system in the PEWE EVAP-WL-161 Cell."

The second and third paragraphs of Section D-1 of the Part B Permit Application under the heading, "VES-WL-135, VES-WL-136, VES-WL-137, VES-WL-138, VES-WL-139, VES-WL-142, VES-WL-144, and VES-WL-150, Process Waste Liquid (PWL) Tanks and Sumps" were revised to read:

"The PWL tanks and sumps are located in CPP-604, CPP-649, and associated valve boxes. The purpose of the system is to collect offgas condensate and liquid from floor drains and transfer the waste to the PEWE Evaporator Feed Collection Tank, VES-WL-133. The system is comprised of tanks VES-WL-135, -136, -137, -138, -139, -142, -144, -150, and various cell sumps. VES-WL-150 collects liquids from the floor drains and the other tanks collect offgas condensate. A sump or vault secondarily contains each of the tanks.

"Sumps SU-WL-140, -143, -145, -146, -147 and -148 do not contain PWL tanks. Sumps SU-WL-140 and SU-WL-143 are located in the Rare Gas Plant in CPP-604. Since the Rare Gas Plant is no longer active there are no sources of waste that would be collected in either sump. Sump SU-WL-148 is located at the INTEC main stack. In the event of equipment failure, condensate from the main stack could collect in this sump. These sumps are emergency equipment and do not routinely collect waste; therefore, the sumps

are exempt from requiring secondary containment. Sumps SU-WL-145, SU-WL-146, and SU-WL-147 are part of the secondary containment and leak detection system for the PEWE Condensate Collection Cell and PEWE EVAP-WL-161 Cell. As such, these sumps do not require the emergency structure exemption set forth in IDAPA 58.01.05.008 [40 CFR § 264.1(g)(8)(i)]."

23. Include the response to NOD #72 in the revised Section C to justify the transfer of PEWE bottoms to the ETS.

#### **RESPONSE:**

The following description was added at the end of the second paragraph in Section D-1 of the Part B Permit Application under the heading, "PEWE System Operation":

"All of these tanks were designed and constructed to contain the types of solutions stored. The P.E. certifications for these units attest that the tank systems are adequately designed and are compatible with the waste(s) to be stored or treated in accordance with IDAPA 58.01.05.008 [40 CFR § 264.192(a)].

"As described above, there may be instances where complete concentration of the waste feed does not occur. When this happens, the remaining feed may be blended with other wastes and reintroduced to the ILWMS. Depending on the characteristics of the new feed solution (e.g., high chlorides, fluorides, or radionuclide concentration), it may be appropriate to route the mixture back to the ETS for processing, rather than the PEWE, to ensure optimum treatment and protection of equipment.

"With the addition of the C-40 valve box, the PEWE bottoms (from both VES-WL-101 and VES-WL-111) can be transferred to the ETS, TFF, and the CPP-604 TFT. From the CPP-604 TFT, waste can be transferred to the TFF, the ETS, or the PEWE. The transfer lines are encased in stainless steel and equipped with leak detection. Drawings showing transfer routes are included in the Section D Plant Drawing package for the Part B Permit Application, Appendix D-1."

Include the response to NOD #74 in the revised Section C to justify the transfer of PEWE process condensate to the LET&D facility or back to the evaporator feed tank.

#### **RESPONSE:**

The following description was added after the second paragraph of Section D-1 of the Part B Permit Application under the heading, "PEWE System Operation":

"Occasionally, PEWE process condensate does not meet the feed limits or operational constraints (e.g., fluorides, TOC, radionuclide concentration) established for the LET&D facility, as identified in Section D-8b(5) of the Part B Permit Application. In these instances, the condensate is routed back to the evaporator feed tank and blended with other solutions for further processing."

#### D-2. Tank Systems

# D-2d. Description of Feed Systems, Safety Cutoffs, Bypass Systems, and Pressure Control: IDAPA 58.01.05.012 [40 CFR § 270.16(c)]

#### **PEWE System**

25. Include the response to NOD #86 in the revised Section D.

#### **RESPONSE:**

The following description was added after the first paragraph of Section D-2d of the Part B Permit Application under the heading, "PEWE Safety Cutoffs":

"The DCS monitors and controls processes in the ILWMS. These processes include the LET&D, Service Waste, PEWE, Process Offgas (POG), Atmospheric Protection System (APS), PWL, VOG, and Main Stack Monitor processes or systems. The DCS is a microprocessor-based control system that uses a combination of free-standing operator consoles networked to electronic I/O interfaces to field devices.

"To ensure a high degree of integrity, redundancy is used where possible. These include redundant controllers, power supplies, communications modules, consoles, and data highway cabling. This redundancy, along with utilization of equipment only from a vendor with documented previous experience of providing successful complex process control systems, and adherence to the vendor's recommended preventive maintenance practices provide the necessary assurance of reliability for meeting the requirements of EPA regulations, Technical Specifications/Standards, and plant mission."

26. Response to NOD #93 does not address DEQ's stated concern. Revise Section D to include a detailed justification for the direct transfer of waste through the evaporators to the PEWE Bottom tanks without operating the evaporators.

#### **RESPONSE:**

Section D-2d of the Part B Permit Application under the heading, "PEWE Bypass Systems" was revised to read:

"Waste can be transferred through the evaporators to VES-WL-101 or VES-WL-111 without operating the evaporators. Such transfers may occur when:

- The evaporators are not operable due to scheduled maintenance activities or are in need of repairs
- The system requires testing following repairs or maintenance (this minimizes waste by not introducing new materials to the system)
- The PEWE Feed Tanks can be emptied to allow additional storage capacity during periods of system maintenance and/or testing."

27. Include the response to NOD #94 to the revised Section D. At a minimum, reference Section D-8b, which describes the CPP-604 building ventilation system.

#### **RESPONSE:**

The first paragraph of Section D-2d of the Part B Permit Application under the heading, "PEWE Pressure Controls" was revised to read:

"Waste treatment and storage vessels at INTEC are connected to a gaseous waste treatment system called the VOG system. All ILWMS storage and treatment systems discharge gases such as instrument air purges (used in level, density, and pressure instrumentation), air spargers (agitators), and gases displaced from a vessel when it fills with liquid.

"Gases from the PEWE and other INTEC processes, such as the Tank Farm and CPP-659 vessels, vent to the CPP-604 VOG system. The CPP-604 VOG system consists of a mist eliminator, superheater, and HEPA filter banks. The mist eliminator and superheater are included to protect and extend the life expectancy of the HEPA filters.

"Several facility process and vessel offgas systems, including the CPP-604 VOG system, combine in the Process APS located in CPP-649. The Process APS is a back-up system that treats the combined process and vessel offgas streams from CPP-601, CPP-604, Tank Farm, and the NWCF. The Process APS treatment consists of a mist eliminator, superheater, and HEPA filters. Next, the process and vessel offgases are routed to the INTEC Main Stack (CPP-708) where they mix with building ventilation air and are exhausted to the atmosphere. The vessel and process offgas systems are maintained under a vacuum to control contamination. The system equipment and piping are fabricated from acid resistant stainless steel for corrosion resistance. Additionally, the Process Condensate Collection Tanks can be vented to the process condensate collection cells, which vent to the CPP-604 building ventilation system. The VOG and APS systems are described further in Section D-8b of this application."

# D-2f(1)(b). Requirements for Secondary Containment and Leak Detection: IDAPA 58.01.05.008 and .012 [40 CFR §§ 264.193 and 270.16(g)]

28. Include the response to NOD #112 to the revised Section D providing details to demonstrate how spilled or leaked waste and accumulated precipitation will be removed from all sumps within the timeframe in accordance with IDAPA 58.01.05.008 [40 CFR § 264.193(b)(3)].

#### **RESPONSE:**

The following description was added after the fourth paragraph of Section D-2f(1)(b) of the Part B Permit Application:

"Upon detection of spilled or leaked materials, the following actions are taken:

 Within 24 hours, remove as much of the waste as is necessary to prevent further releases of hazardous waste to the environment and to allow inspection and repair of the treatment system, in accordance with IDAPA 58.01.05.008 [40 CFR § 264.601]

- Prevent migration of and remove visible contamination from soil or surface water, in accordance with IDAPA 58.01.05.008 [40 CFR § 264.601]
- If the collected material is an HWMA/RCRA-regulated material, manage it in accordance with all applicable requirements of IDAPA 58.01.05.005 through 58.01.05.008 [40 CFR Parts 261 through 264]."
- 29. Include the response to NOD #116 to the revised Section D.

#### **RESPONSE:**

The first paragraph of Section D-2f(1)(b) of the Part B Permit Application under the heading, "CPP-601 Deep Tanks Vaults" was revised to read:

"The WG/WH tanks are located on the lowest level of CPP-601. Two tanks are located in each of the two reinforced concrete vaults. The cells each measure 38 ft 6 in. by 15 ft by 21 ft 6 in., with a stainless-steel-lined floor that extends 3-ft up the walls. Both vaults are provided with sumps and leak detection. Upon high level alarm, the sumps are jetted back to either VES-WG-100/-101 or VES-WH-100/-101 tanks."

30. The revised Part B Permit Application must provide measures for compliance with the applicable secondary containment and leak detection requirements for sections of piping embedded in concrete. To address these requirements, DOE may provide a definitive schedule for completion of any necessary upgrades associated with these embedded lines.

#### **RESPONSE:**

The following concrete penetrations at the INTEC have been identified as requiring upgrades to provide compatible secondary containment per DEQ's guidance:

Penetration Number	Location	Function
3" PWM-1018Y	VES-WM-101/102 Vault	Jet transfer line from WM-101 to WM-100
3" PWM-10024Y	VES-WM-101/102 Vault	Jet transfer line from WM-100 to WM-102
3" PWM-20015Y	VES-WM-101/102 Vault	Overfill line between WM-100/WM-101
1½" PWL-2091C <sup>a</sup>	Wall between 161 Evaporator Cell and the Condensate Collection Cell	Transfer line from VES-WL-109 to the VES-WL-161
1½" PWL-2091C <sup>a</sup>	Wall between 161 Evaporator Cell and the Condensate Collection Cell	Transfer line from VES-WL-109 to the VES-WL-161
4" PWL-1133C	Wall between 161 Evaporator Cell and VES-WL-101 Vault	Evaporators discharge line to VES-WL-101 Tank

Penetration Number	Location	Function
2"PWL-2068C	Condensate Collection Cell to Operating Corridor CPP-604	Discharge line from the collection tanks to the LET&D Process
1" PSAR- 107694 <sup>b</sup>	Wall between the Service Corridor and E Cell.	Drain line from VES-E-108 to PEW
1 ½ " PE-AR-151820 <sup>b</sup>	Wall between the Service Corridor and C Cell. Drawing # 094762/ 133610	Drain line from VES-C-103 to PEW
<sup>3</sup> / <sub>4</sub> " TC-2091Y <sup>b</sup>	Floor from the PM Deck of CPP-601 to the VES-C-103 Drawing # 133610	Drain line from the HOT Sink on the PM Deck to the collection tank VES-C103

a These penetrations do not require core drilling to install a secondary containment sleeve. Adequate area exists around the transfer line to properly sleeve the penetration.

The second and third paragraphs and Table D-5 of Section D-2f(1)(b) under the heading, "VES-WM-101/VES-WM-102 Vault" were deleted and replaced with the following:

"Concrete-embedded transfer lines have been identified at the ILWMS. In order to ensure compliance with the requirements of 40 CFR § 264.193(f), these lines will be upgraded or rerouted in accordance with the following schedule:

- Conceptual design complete by 9/30/04
- Title design complete by 9/30/05
- Work package development complete by 3/31/06
- Identified lines upgraded/rerouted and Professional Engineer certifications submitted to the DEQ on or before 9/30/06."

#### F. PROCEDURES TO PREVENT HAZARDS

# F-2a. General Inspection Requirements: IDAPA 58.01.05.008 and .012 [40 CFR §§ 264.15(a) and (b), 264.33, 264.195, and 270.14(b)(5)]

31. Include the response to NOD #132 to the revised Part B Permit Application, or replace Footnote (1) in Appendix F1-24 with a brief summary of the response to NOD #132.

#### **RESPONSE:**

The following description was added to Section D-2f(1)(b) of the Part B Permit Application under the heading, "EVAP-WL-161 Cell":

b These penetrations will be rerouted to compliant lines as part of a Voluntary Consent Order Action Plan. The work plan is complete awaiting approval of an updated Safety Analysis Report.

"There are known defects (cracks) in the CPP-604 EVAP-WL-161 Evaporator cell concrete walls. INTEC structural engineering personnel have evaluated the condition of the WL-161 Evaporator cell and have determined the cell to be structurally sound. RCRA regulations [40 CFR § 264.15(c)] require repair of structures to ensure the problem does not lead to an environmental or human health hazard. The existing condition of the WL-161 cell does not pose a hazard to the environment or to human health. The portion of the cell that provides secondary containment and leak detection is the lower three-foot stainless steel cell liner, which has no defects. The defects in the cell are only in portions of the concrete walls located above the cell liner. Any leaks from the process vessels or ancillary piping will be completely contained within the stainless steel liner and will not subject the concrete wall to any sustained exposure to hazardous waste. The cell is maintained under negative pressure. Any offgas from a leak or spill would be collected in the CPP-604 VOG system. Inspections of the cell during periods of maintenance or repair are made to ensure that deterioration of the concrete does not increase."

#### F-3a(1). Equipment Requirements: IDAPA 58.01.05.008 and .012 [40 CFR §§ 264.32 and 270.14(b)]

32. Include the response to NOD #133 to the revised Section F.

#### **RESPONSE:**

Appendices F-2 and F-5 in the Part B Permit Application identify the inspection schedules for the PEW evaporators and the LET&D fractionators, respectively.

Form INTEC-4005, "RCRA PEW Tank Overfill And Daily Leak Inspection," is included in Section F, Appendix 1 of the application. This form shows the types of inspections completed for the evaporators.

Form INTEC-4055, "RCRA LET&D Daily Facility Inspections," is included in Section F, Appendix 1 of the application. This form shows the types of inspections completed for the fractionators.

Section F-2a(1) of the Part B Permit Application was revised to read:

"The inspection schedules for the units that comprise the ILWMS, including the scope of the inspections performed and the types of problems noted, are summarized in Appendices F-2 through F-6."

#### F-4. Preventive Procedures, Structures, and Equipment

33. Include the response to NOD #136 to the revised Section F.

#### **RESPONSE:**

The first paragraph of Section F-4a of the Part B Permit Application under the heading, "Unloading Operations" was revised to read:

"Transfers of hazardous waste to and from CPP-604 are conducted through piping systems. Wastes generated at other INEEL or off-Site facilities may be introduced to the ILWMS via the CPP-1619 Truck Unloading Bay through tanker trucks or containers.

Unloading operations at this facility are controlled by standard operating procedures. During unloading operations, a portable HEPA air mover is required to filter particulate and radioactive emissions. An intake for the air mover is located near the unloading hose connections in the CPP-1619 unloading bay. A stainless-steel drip pan is used to collect possible leaks during unloading. Personnel will inspect for evidence of improper connections before beginning the transfer or acceptance of waste at CPP-1619. Waste staging areas will be inspected for leaks or spills when waste is being received."

#### F-4d. Equipment and Power Failure: IDAPA 58.01.05.012 [40 CFR § 270.14(b)(8)(iv)]

34. Include the response to NOD #139 to the revised Section F.

#### **RESPONSE:**

Section F-4d of the Part B Permit Application was revised to read:

"Some components of the ILWMS are supplied with redundant equipment. If equipment should fail on these systems, it would have minimal effect on the operating unit, since the redundant equipment would be started and the operation stabilized. The failed equipment would then be investigated to determine the cause of the failure, and repairs would be initiated. If a system that did not have redundant equipment were to fail, the operating unit would be secured.

"Upon total loss of electrical power, ILWMS equipment that manages hazardous and mixed wastes is designed to shut down in a manner that protects employees, equipment, human health, and the environment.

"Cranes and hoists are considered non-critical equipment and are not supplied with emergency standby power. This type of equipment is designed to fail in place. Movement will be suspended until power is restored.

"The DCS is designed with battery backup to maintain operability and to ensure safe shutdown.

### **CPP-604 PEWE System and TFT**

"The Evaporator Feed Collection Tank (VES-WL-133), the Process Condensate Surge Tank (VES-WL-131) and the Process Condensate Collection Tanks (VES-WL-106, VES-WL-107, and VES-WL-163) are all equipped with two redundant transfer pumps.

"The PEW evaporators (EVAP-WL-129 and EVAP-WL-161) and associated heat exchangers are identical and may be operated independently or in parallel.

"PEW evaporator bottoms can be stored/treated in either VES-WL-101 or VES-WL-111.

#### **CPP-601 Deep Tanks**

"There are a total of four Deep Tanks (VES-WG-100, VES-WG-101, VES-WH-100, and VES-WH-101) that are essentially redundant systems. Each tank is equipped with sparge flow instruments, level instrumentation, one transfer jet, and one transfer pump.

"Sufficient redundancy exists such that a receiving tank is available for collection. Loss of a sparge flow or level instrument may require an operator to switch collection to another tank, but would not require total cessation of operations.

"If a system that does not have redundant equipment was to fail, the operating process would be shut down and not operated again until the failure was repaired.

#### **CPP-1618 LET&D Facility**

"The LET&D fractionators (FRAC-WLL-170 and FRAC-WLK-171) and associated heat exchangers are identical.

"The Acid Fractionator Bottoms Tank (VES-WLL-195) is equipped with two redundant transfer pumps."

35. Include the response to NOD #141 to the revised Section F.

#### **RESPONSE:**

See the response to item No. 34 of this NOD.

#### F-4f. Releases to the Atmosphere: IDAPA 58.01.05.012 [40 CFR § 270.14(b)(8)(iv)]

36. Since the CPP-641 Westside Waste Holdup tanks have been emptied to the maximum extent allowed, the current WWH tank vault configuration, although they are not connected to the VOG system, is acceptable. When/if the WWH tank system is upgraded, vault offgas system improvements must be taken into consideration.

#### **RESPONSE:**

The INEEL concurs. When/if the WWH tank system is upgraded, those upgrades would include replacement of the tile-encased transfer lines with piping that has secondary containment compatible with corrosives, lining of the vaults with stainless steel, an upgrade of the instrumentation, a sampler upgrade, and vessel/vault offgas system improvements.

#### I. CLOSURE AND POST-CLOSURE REQUIREMENTS

37. Include the response to NOD #151 and #152 to the revised Section I.

#### **RESPONSE:**

The following description was added after first paragraph of Section I 1.1.2(a) of the Part B Permit Application:

"The closure plan will be modified, in accordance with IDAPA 58.01.05.008 [40 CFR § 264.112(c)], to include the appropriate verification sampling techniques to be used to meet the closure performance standards prior to implementation of the closure plan."

The same description was added to Section I 1.1.3(a)(4) of the Part B Permit Application.

#### **Other Changes**

- 1. The second paragraph of Section C-1g of the Part B Permit Application was revised to state that PEWE bottoms may be stored in compliant storage units, other than the Tank Farm Facility (TFF), until an ultimate treatment and disposal alternative has been selected, permitted, and constructed. In an effort to empty and close TFF tanks, wastes may be transferred from the TFF to other RCRA-compliant storage units prior to permitting and construction of the ultimate treatment alternative for evaporator bottoms.
- 2. The bulleted item, immiscible organic liquids, was removed from the list of prohibited items in Section C-2a(1) of the Part B Permit Application. Small quantities of immiscible organics may be managed by the ILWMS as long as the system process tolerance limits are met.
- 3. Two parameters, specific gravity and total inorganic carbon, were removed from Table C-3 of the Part B Permit Application as parameters that are evaluated for process samples taken from the CPP-601 Deep Tanks. These parameters were inadvertently included in the table, which was added in response to the first NOD for this application. Although these parameters are evaluated for characterization samples, they are not analyzed for process samples.
- 4. Examples of inspection forms provided in Appendix F-1 of the Part B Permit Application were updated to reflect the most current revision of each form.



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## **Material Safety Data Sheet**

Material Name: ALKALINE RUST REMOVER T-4181

ID: 238022

## \* \* Section 1 - Chemical Product and Company Identification \*

#### Product Trade Name ALKALINE RUST REMOVER T-4181

Manufacturer Information

Henkel Surface Technologies
Henkel Corporation
32100 Stantaneon Highway

(248) 583-9300

32100 Stephenson Highway Madison Heights, MI 48071

(800) 424-9300

# \* \* \* Section 2 - Composition / Information on Ingredients \* \* \*

CAS#	Component	
1310-73-2	Sodium hydroxide	Percent
102-71-6		> 60
	Triethanolamine	10-20
527-07-1	Sodium gluconate	1-10
111-42-2	Diethanolamine	1-10

## \* Section 3 - Hazards Identification \* \* \*

#### **Emergency Overview:**

DANGER!

CAUSES EYE AND SKIN BURNS. MAY CAUSE BLINDNESS.

CAUSES DIGESTIVE TRACT BURNS.

REPEATED OR PROLONGED EXPOSURE MAY CAUSE LIVER AND KIDNEY DAMAGE BASED ON ANIMAL DATA

EVEN DILUTE SOLUTIONS MAY CAUSE BURNS.

#### Potential Health Effects:

Inhalation and skin contact are expected to be the primary routes of occupational exposure to this material. Based on its composition, it is anticipated to be corrosive to the eyes, skin and respiratory tract. Inhalation of mist or vapor may cause coughing, sore throat, shortness of breath, lung injury or chemical pneumonia. Studies with animals repeatedly exposed to components of this material have produced liver and kidney damage. Medical conditions which may be aggravated by exposure to this material include lung, liver or kidney disease or limited respiratory capacity.

### \* \* Section 4 - First Aid Measures \* \* \*

#### Eye Contact:

#### **\$kin Contact:**

Immediately flush with plenty of water for at least 30 minutes while removing contaminated clothing and shoes. Get medical attention immediately. Wash clothing before reuse. Destroy contaminated shoes.

#### Ingestion:

Do NOT induce vomiting. Give water to drink. Get medical attention immediately. NEVER GIVE ANYTHING BY MOUTH TO AN UNCONSCIOUS PERSON.

#### Inhalation:

Remove to fresh air. If not breathing, give artificial respiration. If breathing is difficult, give oxygen. Get medical attention.

# \* \* \* Section 5 - Fire Fighting Measures \* \* \*

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Material Name: ALKALINE RUST REMOVER T-4181

ID: 238022

Flash Point: NA

Method Used:

Flammability Classification:

Upper Flammable NA

Lower Flammable NA Limit (LFL):

Limit (UFL):

Fire & Explosion Hazards:

Avoid breathing fumes from fire exposed material.

Extinguishing Media:

Use water spray or water fog

Fire-Fighting Instructions:

Use water spray to cool containers exposed to fire. Fire fighters and others who may be exposed to products of combustion should wear full fire fighting turn out gear (full Bunker Gear) and self-contained breathing apparatus (pressure demand NIOSH approved or equivalent). Fire fighting equipment should be thoroughly decontaminated after use.

## \* \* \* Section 6 - Accidental Release Measures \* \* \*

#### Spill or Leak

Stop the leak, if possible. Ventilate the space involved. Contain, sweep up, place in container for disposal. Shut off or remove all ignition sources. Prevent waterway contamination. Construct a dike to prevent spreading. Collect run-off and transfer to drums or tanks for later disposal.

Clean up procedures: Transfer to containers in preparation for later disposal. Avoid generation of vapors. Place in non-sparking containers for recovery or disposal. Remove from spill location. Decontaminate area.

## \* \* \* Section 7 - Handling and Storage \* \* \*

#### Handling Procedures:

Do not breathe dust. Do not get in eyes, on skin or clothing. Wash thoroughly after handling. Keep container tightly closed. Empty container may contain hazardous residues. Do not enter confined spaces unless adequately ventilated. To avoid rapid temperature rise, violent spattering, or explosive eruptions: always add caustic to water when mixing. Never add water to a caustic when mixing.

#### Storage Procedures:

Store in a cool, dry place. Avoid excessive heat. Store out of direct sunlight in a cool, well-ventilated place.

# \*\*\* Section 8 - Exposure Controls / Personal Protection \*\*\*

#### Component Exposure Limits

Sodium hydroxide (1310-73-2)

ACGIH: C 2 mg/m3 OSHA: 2 mg/m3 TWA NIOSH: C 2 mg/m3

Triethanolamine (102-71-6)

ACGIH: 5 mg/m3 TWA

Diethanolamine (111-42-2)

ACGIH: 2 mg/m3 TWA

skin - potential for cutaneous absorption

NIOSH: 3 ppm TWA; 15 mg/m3 TWA

#### Engineering Controls:

Investigate engineering techniques to reduce exposures below airborne exposure limits. Provide ventilation if necessary to control exposure levels below airborne exposure limits (see below). If practical, use local mechanical exhaust ventilation at sources of air contamination such as open process equipment. Consult ACGIH ventilation manual or NFPA Standard 91 for design of exhaust systems.

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Material Name: ALKALINE RUST REMOVER T-4181

ID: 238022

#### PERSONAL PROTECTIVE EQUIPMENT

As prescribed in the OSHA Standard for Personal Protective Equipment (29 CFR 1910.132), employers must perform a Hazard Assessment of all workplaces to determine the need for, and selection of, proper protective equipment for each task performed.

#### Eyes/Face Protective Equipment:

Where there is potential for eye contact, wear a face shield, chemical goggles, and have eye flushing equipment immediately available.

#### Skin Protection:

Wear appropriate chemical resistant protective clothing and protective gloves to prevent contact. Consult glove manufacturer to determine appropriate type glove material for given application. Rinse immediately if skin is contaminated. Wash contaminated clothing and clean protective equipment before reuse. Provide a safety shower at any location where skin contact can occur. Wash skin thoroughly after handling.

#### Respiratory Protection:

Avoid breathing dust. When airborne exposure limits are exceeded (see below), use NIOSH approved respiratory protection equipment appropriate to the material and/or its components. Consult respirator manufacturer to determine appropriate type equipment for given application. Observe respirator use limitations specified by NIOSH or the manufacturer. For emergency and other conditions where exposure limit may be significantly exceeded, use an approved full face positive-pressure, self-contained breathing apparatus or positive-pressure airline with auxiliary self-contained air supply. Respiratory protection programs must comply with 29 CFR § 1910.134.

## \*\*\* Section 9 - Physical & Chemical Properties \*\*\*

Physical State: Solid

Odor:

Vapor Density: NA Melting Point: NE

pH: > 12.0 VOC: 1.5% (Calculated) Appearance:

Off white, granular mixture

Vapor Pressure: NA Boiling Point: NA

Boiling Point: NA Specific Gravity: NA

Viscosity: Solubility Water: Soluble

## \*\*\* Section 10 - Chemical Stability & Reactivity Information \*\*

#### Chemical Stability:

This material is chemically stable under normal and anticipated storage and handling conditions.

#### Incompatibility:

Avoid contact with strong acids. Contains organic amine compounds. Nitrite based materials should not be added due to possible nitrosoamine formation.

#### **Decomposition Products:**

Oxides of carbon, nitrogen compounds.

#### Hazardous Polymerization:

Hazardous polymerization is not known to occur.

## \* \* \* Section 11 - Toxicological Information \* \* \*

#### Acute Toxicity:

#### A: General Product Information

No information available for the product.

#### B: Component Analysis - LD50/LC50

Triethanolamine (102-71-6)
Oral LD50 Rat: 4920 uL/kg
Oral LD50 Mouse: 5846 mg/kg
Dermal LD50 Rabbit: >20 mL/kg

Diethanolamine (111-42-2)

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· Make one recognized

Material Name: ALKALINE RUST REMOVER T-4181

ID: 238022

Oral LD50 Rat: 620 uL/kg Oral LD50 Mouse: 3300 mg/kg Dermal LD50 Rabbit: 7640 uL/kg

#### Component Carcinogenicity

Diethanolamine (111-42-2)

IARC: Monograph 77, 2000 (Group 3 (not classifiable))

#### **Chronic Toxicity**

No information available for the product.

#### **Epidemiology:**

No information available for the product.

#### **Neurotoxicity:**

No information available for the product.

#### Mutagenicity:

No information available for the product.

#### Teratogenicity:

No information available for the product.

## Section 12 - Ecological Information

#### **Ecotoxicity:**

## A: General Product information

No information available for the product.

## **B: Component Analysis - Ecotoxicity - Aquatic Toxicity**

Triethanolamine (102-71-6)

**Test & Species** 

Conditions

LC50 (24 hr)

5000 mg/L.

goldfish

Diethanolamine (111-42-2)

**Test & Species** 

Conditions

LC50 (96 hr) fathead minnow LC50 (24 hr) goldfish

LC50 (24 hr) goldfish

>100 mg/L. 800 mg/L. 5000 mg/L.

Static. pH 9.6. pH 7.0. 15 °C.

EC50 (5 min) Photobacterium phosphoreum

73 mg/L Microtox

#### Environmental Fate:

No data is available concerning the environmental fate, biodegradation or bioconcentration for this product.

## \* \* \* Section 13 - Disposal Considerations

## **US EPA Waste Numbers & Descriptions:**

#### A: General Product Information

Recover, reclaim or recycle when practical. Dispose of in accordance with federal, state and local regulations. Note: Chemical additions to, processing of, or otherwise altering this material may make this waste management information incomplete, inaccurate, or otherwise inappropriate. Furthermore, state and local waste disposal requirements may be more restrictive or otherwise different from federal laws and regulations.

#### **B: Component Waste Numbers**

No EPA Waste Numbers are applicable for this product's components.

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## **Material Safety Data Sheet**

Material Name: ALKALINE RUST REMOVER T-4181

ID: 238022

## Section 14 - Transportation Information \* \* \*

**US DOT Information** 

Shipping Name: Please refer to the container label for transportation information.

\* \* \* Section 15 - Regulatory Information \* \* \*

### **US Federal Regulations**

### A: General Product Information

This product is considered hazardous under 29 CFR 1910.1200 (Hazard Communication).

#### **B: Component Analysis**

This material contains one or more of the following chemicals required to be identified under SARA Section 302 (40 CFR 355 Appendix A), SARA Section 313 (40 CFR 372.65) and/or CERCLA (40 CFR 302.4).

Sodium hydroxide (1310-73-2)

CERCLA: final RQ = 1000 pounds (454 kg)

#### Diethanolamine (111-42-2)

SARA 313: form R reporting required for 1.0% de minimis concentration CERCLA: final RQ = 100 pounds (45.4 kg)

SARA 311/312: Acute: Y Chronic: Y Fire: N Pressure: N Reactive: N

#### State Regulations

### A: General Product Information

No additional information available.

#### B: Component Analysis - State

The following components appear on one or more of the following state hazardous substances lists:

Сотролелт	CAS#	CA	I FL	MA	MN	TNJ	PA
Sodium hydroxide	1310-73-2	Yes	Yes	Yes	Yes	Yes	Yes
Triethanolamine	102-71-6	No	Yes	Yes	Yes	No	Yes
Diethanolamine	111-42-2	Yes	Yes	Yes	Yes	Yes	Yes

## Other Regulations

## A: General Product Information

All components are on the U.S. EPA TSCA Inventory List.

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Material Name: ALKALINE RUST REMOVER T-4181

ID: 238022

## B: Component Analysis - InventoryComponent Analysis - Inventory

Component	CAS#	TSCA	DSL	EINECS
Sodium hydroxide	1310-73-2	Yes	Yes	Yes
Triethanolamine	102-71-6	Yes	Yes	Yes
Diethanolamine	111-42-2	Yes	Yes	Yes
Sodium gluconate	527-07-1	Yes	Yes	Yes

#### C: Component Analysis - WHMIS IDL

The following components are identified under the Canadian Hazardous Products Act Ingredient Disclosure List:

Component	CAS#	S Products Act Ingredient Disclos  Minimum Concentration
Sodium hydroxide	1310-73-2	1%; English Item 1442; French Item 998
Triethanolamine	102-71-6	1%; English Item 1621; French Item 1663
Diethanolamine	111-42-2	1%; English Item 569; French

## \* \* \* Section 16 - Other Information \* \* \*

#### Key/Legend

EPA = Environmental Protection Agency; TSCA = Toxic Substance Control Act; ACGIH = American Conference of Governmental Industrial Hygienists; IARC = International Agency for Research on Cancer; NIOSH = National Institute for Occupational Safety and Health; NTP = National Toxicology Program; OSHA = Occupational Safety and Health Administration; NFPA = National Fire Protection Association; HMIS = Hazardous Material Identification System; CERCLA = Comprehensive Environmental Response, Compensation and Liability Act; SARA = Superfund Amendments and Reauthorization Act

The information presented herein is believed to be factual as it has been derived from the works and opinions of persons believed to be qualified experts; however, nothing contained in this information is to be taken as a warranty or representation for which Henkel Surface Technologies bears legal responsibility. The user should review any recommendations in the specific context of the intended use to determine whether they are appropriate.

Contact: Regulatory Affairs and Product Acceptance

Contact Phone: (248) 593-9300

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## MATERIAL SAFETY DATA SHEET

Westco Chemicale, Inc. 11312 Hartland Street North Hollywood, California 91605 (213) 577-0077 (818) 980-1152

EMERGENCY CONTACT:

CALL CHEMTREC : DAY OR NIGHT

(800)424-9300

SUBSTANCE IDENTIFICATION

144-62-7 OXALIC ACID

HMIS

TRADE NAMES/SYNONYMS-

ETHANEDIOIC ACID

HEALTH----1
FLAMMABILITY-1
REACTIVITY---0

CHEMICAL FAMILY----

ORGANIC ACID

PERSONAL PROTECTION-F

MOLECULAR FORMULA----

HOOCCOOH 2H2O

MOLECULAR WT---126.07

COMPONENTS AND CONTAMINANTS

OXALIC ACID - 99.6% INERT SALTS - 0.4%

NK

PHYSICAL DATA

DESCRIPTION-

WHITE CRYSTALLINE POWDER, ODORLESS

NA NA NA

MELTING PT---101 DEG C BOILING PT----186 DEG C

SOLUBLE - 11.7%

DECOMPOSES % VOLATILE-----NA

SPECIFIC GRAVITY--1.653

HOW TO DETECT THIS SUBSTANCE : CHEMICAL ANALYSIS

FIRE AND EXPLOSION DATA

FLASH POINT-

FIRE EXTINGUISHING MTRLS-

NA

WATER, DRY CHEMICALS, FOAM, CO2

FLAMMABLE LIMITS---NK

SPECIAL FIRE FIGHTING

PROCEDURES -

SELF CONTAINED BREATHING APPARATUS. WEAR FACE SHIELD OR SAFETY GOGGLER

UNUSUAL FIRE AND

EXPLOSION HAZARDS--

NONE, DECOMPOSITION PRODUCTS INCLUDE CARBON

MONOXIDE & FORMIC ACID WHICH ARE TOXIC & FLAMMABLE

## TRANSPORTATION DATA

DOT CLASSIFICATION ----NONE DOT LABEL NONE UNNA NUMBER-----NONE SPECIAL INSTRUCTIONS ----NONE

TOXICITY

DANGER-CORROSIVE

LD/50 - 71 mg/kg 1mg/M3 ACGIH - PEL DO NOT EAT OR SMOKE IN WORK AREA

HEALTH EFFECTS AND FIRST AID

INHALATION

ACUTE EXPOSURE-

CHRONIC EXPOSURE ....

FIRST AID

TOXIC-INHALATION OF DUST IS POISONOUS

REMOVE TO FRESH AIR. ADMINISTER OXYGEN IF NECESSARY

SKIN CONTACT

ACUTE EXPOSURE-

STRONG IRRITANT TO SKIN & MUCOUS MEMBRANES. MAY

**CAUSE BURNS** 

CHRONIC EXPOSURE-

FIRST AID----

WASH WITH SOAP & WATER

EYE CONTACT

**ACUTE EXPOSURE-**

CHRONIC EXPOSURE-

FIRST AID-

MAY CAUSE CORNEAL DAMAGE

NK

NK

FLUSH WITH WATER FOR 15 MINUTES. OBTAIN ASSIS-

TANCE FROM OPHI HAMOLOGIST

NGESTION

ACUTE EXPOSURE....

CHRONIC EXPOSURE----

FIRST AID-

POISONOUS IF SWALLOWED. MAY CAUSE GASTROENTERITIS

NK

DO NOT INDUCE VOMITING. GIVE MILK. CALL A DOCTOR

ANTIDOTE ----

MILK OF MAGNESIA OR ANTACIDS

SUSPECTED CANCER AGENT

NO

RECOMMENDATIONS TO A DOCTOR

TREAT SYMPTOMATICALLY. NEUTRALIZE WITH SODIUM

BICARBONATE

#### REACTIVITY DATA

STABILITY .... INCOMPATIBILITIES-DECOMPOSITION PRODUCTS-

STRONG OXIDIZERS

STABLE

HAZARDOUS POLYMERIZATION-CONDITIONS TO AVOID-

FORMIC ACID, CARBON DIOXIDE

WILL NOT OCCUR

HEAT, MOISTURE (HYGROSCOPIC), DUSTING

# SPILL AND LEAK PROCEDURES

OCCUPATIONAL SPILL

SWEEP UP AND CONTAINERIZE, DISSOLVE IN LARGE VOLUME OF WATER AND NEUTRALIZE WITH SODA ASH

WASTE DISPOSAL METHODS-

INCINERATION OR FLUSH INTO SEWER WITH LARGE VOLUME

OF WATER AFTER NEUTRALIZATION

NOTE : DISPOSE OF ALL WASTES IN ACCORDANCE WITH FEDERAL, STATE AND LOCAL

# SPECIAL HANDLING INFORMATION

VENTILATION ...

LOCAL EXHAUST

RESPIRATORY PROTECTION— CLOTHING.

DUST MASKNIOSH RESPIRATOR RUBBER BOOTS, RUBBER APRON

GLOVES----

ALBEER

EYE PROTECTION ----

SAFETY GOGGLES OR FULL FACE MASK

# ADDITIONAL INFORMATION

THIS MSDS WAS OBTAINED FROM RELIABLE SOURCES. HOWEVER, IT IS PROVIDED WITH OUT REPRESENTATION OR WARRANTY EXPRESSED OR MPLIED REGARDING ACCURACY OR CORRECTNESS. CONDITIONS, METHODS OF HANDLING, STORAGE, USE AND DISPOSAL OF THE PRODUCT ARE BEYOND OUR CONTROL OR BEYOND OUR KNOWLEDGE. FOR THIS AND OTHER REASONS, WE ASSUME NO RESPONSIBILITY AND EXPRESSLY DISCLAIM LIABILITY FOR INJURY, LOSS, DAMAGE OR EXPENSES ARISING FROM THE USE OF THIS PRODUCT.

AUTHORIZED BY WESTCO CHEMICALS, INCORPORATED

CREATION DATE-REVISION DATE

5/25/90 3/6/98

NA=NOT APPLICABLE ND=NOT DETERMINED NK=NOT KNOWN

## 005-100

Material Safety Data Sheet May be used exceedy sale OSHA's Heased Communication Standard, 20 CFR 1910.1200, Standard must be consulted for specific/regularyments.		U.S. Department Occupational Safety or (Non-Mendatory Form Form Approved OMB No. 1218-0072	nd Health Administration	<b></b>	
Rediscussis		Many March special are in Interceptor in Gratia	t paymented. If any how is not t in. The space ment he maked	estado, er 10 No industr Per	
Section I	•				
Atomic Products Corp.		(516) 924-900	R		
Address Abantor, Street Chy State, and 20 Cody 49 Nation Orive		(516) 924-900		<del></del>	
Shirley, NJ 11967		Ome 7722788			
		Signato di Presser festi	7		
Section II Hazardove IngredientsAder	tity information		Yestro	····	
Hammithus Components (Specific Chemical Martin); (		ORNA PIEL ACQU	Care Units	% (species	
Citric Acid	Annual Consider	CAS 77-92-9		2.9	
Octyl Phenol Condensed		CAS 9036-19-5		6.0	
with 8 - 10 moles Ethylene Oxi	de Triton XI	100			
			<del></del>		
Tetrasodium Ethylenediamine		CAS 64-02-8		5.7	
Triacetate		<del></del>	<del></del>		
Benzyldimethyl (2-(2-(P- (1,1,	1.1 tetra-n	ethylhutyll		0.008	
Phenoxy) Ethoxy) Ethyl) Ammonia		,,,,,,,			
Hyamine 1622	-	CAS 121-56-0			
Mineral Water	~~-	1.4. (213)450.	<del></del>	85.39	
ection III Physical/Chemical Characte	ristics				
siling Point	100°C	Specific Gravity (H <sub>Z</sub> O - 1)		1.052	
por Pressure (mm Hg.)	-	Meting Point			
spor Dungity (AIR = 1)	NA NA	Freez Evaporation Rate	ing Point	0.3°C	
	NA	( water - 1)		1.2	
Infinite (complete	ly miscibl				
Blufsh transparent liquid - sl					
ection IV — Fire and Explosion Hazard				<del></del>	
Mn Pert (Methos Used)		Flammable Limits	LEL	UEL	
Greater than 214°F		NA			
Ory powder, foam, carbon diox	de				
ecial Five Fighting Procedures Fire fighters should wear self	f-contained	breathing apparatu	ıs.		
Decomposition products may be	****				
necomposition products may be	LUXIC.				

<del></del>	United		Conditions to Avoid						
		1_			<del></del>				
	State	X	Metal nitra	tes					
competibility (	Mareriels to Avail	d) Ma	tallic surfaces	for pro	longed	time De	riods (P	H 5)	
sardeus Deco	mposition or Bypro	drick.				V			
eservieus elymerication	May Occur	T	Constions to Avoid						
	Will Net Coour	X	None						
ection VI —	Heelth Hazan	d Date							
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February 26, 2002 CCN 30195

Mr. J. T. Case, Director INTEC Waste Program U.S. Department of Energy Idaho Operations Office 850 Energy Drive, MS 1154 Idaho Falls, ID 83401-1563

CONTRACT NO. DE-AC07-99ID13727 - TRANSMITTAL OF NWCF EVAPORATOR TANK SYSTEM 2001 OFFGAS EMISSIONS INVENTORY, INEEL/EXT-02-00198

Dear Mr. Case:

A copy of the external report NWCF Evaporator Tank System 2001 Offgas Emissions Inventory is enclosed. This report provides semi-volatile and volatile organic, metals, particulate, and acid gas emissions data and rates for the New Waste Calciner Facility Evaporator Tank System (NWCF ETS) as measured during operations from May through June 2001. A detailed assessment of the data quality has also been provided.

The emissions inventory for the NWCF ETS was completed to: 1) demonstrate the applicability of standard EPA offgas sampling methods and 2) characterize the constituents of primary concern for this offgas stream. Only minor variations to the EPA methods were required to compensate for radiological exposure and safety concerns and the below-grade sampling port location.

Emissions rates for the measured constituents were compared to rates measured for the NWCF Calciner and scaled to NWCF Calciner Screening Level Risk Assessment parameters to provide comparisons to the EPA hazard quotient and cancer risk guidance. The emissions rates and human health risk values for the NWCF ETS were lower than those for the NWCF Calciner.

The enclosed report fulfills a performance measure in Program Execution Guidance EM-D-33 FY-2002 and internal milestone A7131021 in control account C.1.06.01.07.13 Waste Sampling and Characterization.

Questions regarding the content of this report should be addressed to L. J. Young at 526-3132 or R. D. Boardman at 526-3732.

Sincerely,

J. H. Valentine, Manager of Projects

High Level Waste Program

rmg

Enclosure

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Mr. J. T. Case February 26, 2002 CCN 30195 Page 3

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# **NWCF Evaporator Tank System 2001 Offgas Emissions Inventory**

R. D. Boardman K. M. Lamb L. A. Matejka J. A. Nenni

February 2002

Idaho National Engineering and Environmental Laboratory
High Level Waste Program
Bechtel BWXT Idaho, LLC
Idaho Falls, ID 83415

Prepared for the
U.S. Department of Energy
Assistant Secretary for Environmental Management
Under DOE Idaho Operations Office
Contract DE-AC07-99ID13727

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# **ABSTRACT**

An offgas emissions inventory and liquid stream characterization of the Idaho New Waste Calcining Facility (NWCF) Evaporator Tank System (ETS), formerly known as the High Level Liquid Waste Evaporator (HLLWE), has been completed. The emissions rates of volatile and semi-volatile organic compounds, multiple metals, particulate, and hydrochloric acid (HCl)/Cl<sub>2</sub> were measured in accordance with an approved Quality Assurance Project Plan (QAPjP) and Test Plan that invoked U.S. Environmental Protection Agency (EPA) standard sample collection and analysis procedures. Offgas samples were collected during the start up and at the end of evaporator batches when it was hypothesized the emissions would be at peak rates. Corresponding collection of samples from the evaporator feed, overhead condensate, and bottoms was made at approximately the same time as the emissions inventory to support material balance determinations for the evaporator process. The data indicate that organic compound emissions are slightly higher at the beginning of the batch while metals emissions, including mercury, are slightly higher at the end of the evaporator batch. The maximum emissions concentrations are low for all constituents of primary concern. Mercury emissions were less than 5 ppbv (< 40 μg/dscm), while the sum of HCl and Cl<sub>2</sub> emissions was less than 1 ppmv. The sum of all organic emissions also was less than 1 ppmv. The estimated hazardous quotient (HQ) for the evaporator was 6.2e-6 as compared to 0.25 for the EPA target criteria. The cancer risk was 1.3e-10 compared to an EPA target of 1e-5.

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## **SUMMARY**

This report presents the 2001 effluent gas emissions inventory data for the NWCF Evaporator Tank System (ETS) operated at the INTEC. Liquid wastes generated from decontamination activities are stored in the INTEC High Level Waste Tank Farm Facility (TFF). The Tank Farm wastes are currently being concentrated using the NWCF ETS (formally known as High Level Liquid Waste Evaporator, or HLLWE). The NWCF ETS currently operates under Resource Conservation and Recovery Act (RCRA) interim status. A RCRA Part B permit application for this unit is being prepared and will be submitted in FY-2003. In order to support the permitting activities, the feed and output streams were characterized during evaporator operations in May and June, 2001. During this time, the NWCF ETS was being used to reduce the volume of a blend of two parts by volume of solution from WM-184 and one part by volume of solution from WM-181. Both of these tanks contained sodium-bearing waste (SBW).

The NWCF ETS is a single-stage, thermal siphon, batch evaporator. Dilute Tank Farm liquid wastes are semi-continuously fed to the evaporator to maintain a constant level in the evaporator. The system consists of a feed tank (VES-NCC-152), a flash column (VES-NCC-150), a reboiler (HE-NCC-350), and a condenser (VES-NCC-151). Blended tank farm wastes are added to the flash column via the feed tank. When the level in the flash column reaches its normal operating level, steam is introduced into the shell side of the evaporator reboiler. Once the desired specific gravity is reached, the concentrated solution is cooled and returned to the Tank Farm. Each batch generally requires 10-16 hours to process, followed by 10-16 hours to refill the feed tanks and to attend to waste transfers back to the Tank Farm.

The NWCF ETS overhead vapor is condensed and sent to the INTEC Process Equipment Waste Evaporator (PEWE) to be re-evaporated. Non condensable and purge gasses are vented from the feed tank and the condenser to the NWCF equipment vent system. The combined offgas passes through the NWCF high-efficiency particulate air (HEPA) filters and then through the Atmospheric Protection System (APS) before being discharged from the main INTEC stack with other vessel offgas and building ventilation air. The offgas tie-in sample location previously used to sample the NWCF Calciner offgas stream was determined to be the best location for sampling the NWCF ETS emissions.

#### Scope and Approach

The Tank Farm wastes are highly acidic (mainly nitric acid) and contain several RCRA metals, including mercury, and trace amounts of volatile and semi-volatile hazardous organic compounds which were introduced into the Tank Farm Facility by previous disposal of laboratory analytical wastes, NWCF Calciner scrub solution recycle, and organic solvent cleaning. Those components with low boiling points are released to the NWCF ETS offgas system during waste transfers, mixing, and evaporation. Trace amounts of heavy, nonvolatile hydrocarbons may also exist in the waste feed solutions; however, emissions of the nonvolatile hydrocarbons and also the nonvolatile metals may occur due to aerosol entrainment from the evaporator. The majority of the entrained droplets are believed to be collected in the offgas condensers, mist eliminators, and HEPA filters.

The scope of the NWCF ETS process effluent gas emissions inventory included:

- Measurement of the NWCF offgas duct velocity, temperature, and flowrate during operation of the NWCF ETS
- Manual offgas sampling and analysis for particulate matter (PM), HCl, Cl<sub>2</sub>, selected metals, volatile organic compounds (VOCs), and semi-volatile organic compounds (SVOCs)
- Measurement of oxygen concentrations

Standard EPA sample collection and analysis methods were used to collect the offgas samples. Sampling was conducted following standard EPA methodology for emissions compliance testing, with attention being given to the following:

- Development and adherence to an approved project quality assurance/quality control plan
- Implementation of chain-of-custody (COC)/requests-for-analysis (RFA) and master sample collection lists that utilize and implement an in-field sample tracking and sample identification number verification
- Development of target analyte lists (TAL) and precision, accuracy, representativeness, completeness, and comparability (PARCC) data quality objectives
- Collection of samples using checklists to record train set up, sample collection data, and sample recovery steps
- Collection and analysis of reagent blanks, trip blanks, and field blanks to achieve prescribed data quality objectives
- Sample collection monitoring by a Project Quality Assurance Officer (PQAO)
- Application of EPA Solid Waste (SW)-846 and 40 Code of Federal Regulations (CFR) 60 Appendix A reference methods for sample analysis.
- Multiple reviews and verifications of field data, analytical data, process data, and resultant calculations of emissions rates

Samples of the feed were analyzed prior to initiation of processing the blend in the NWCF ETS to ensure that the chemistry of the feed solution was compatible with the process equipment. Samples of the condensed overheads and the concentrated bottoms from the first several batches processed were analyzed to ensure that the chemistry of those streams was compatible with down-stream process equipment. The results of these analyses have been included in this report to provide a resource for process permitting discussions and planning.

At the beginning of the offgas sample collection period, the vertical gas velocity profile and swirl angle in the duct were measured to determine an appropriate fixed-point location to collect the offgas samples. Sample contamination survey trains and routine

radiological surveys and screenings were completed throughout the sample collection period to ensure that the samples shipped to the contract analytical laboratory met the labs radioactive materials license criteria. At the end of the sample collection period, the sample probe was removed from the duct and rinsed with acetone and nitric acid. The acetone and nitric acid probe rinses were analyzed for PM and metals.

A set of two runs was completed for each EPA sample train configuration at the beginning and another at the end of evaporator batches. This provided a total of four runs for each method to compare emissions trends at the beginning and end of the evaporator batches. Oxygen concentrations were monitored during each sample train run. The oxygen concentrations in the duct were consistently found to be similar to ambient air conditions. Therefore, it was not necessary to constantly operate the oxygen monitor.

Moisture levels in the offgas were determined from gravimetric and/or volumetric changes in the sample train resins and impingers, respectively. The offgas moisture level was typically less than the dew point of the sample gas passing through the sample collection train condensers. At the most, only 1-2 mL net condensate was collected in any of the condensate knockout impingers.

## Data Quality Assessment

All of the planned emissions inventory samples data and associated quality assurance/quality control (QA/QC) samples were collected in accordance with the test plan (PLN-879) and Quality Assurance Project Plan (PLN-880) which were developed and approved for this project. An extensive discussion is provided in the report body regarding conformance of the sample collection activities with the procedures and EPA Method requirements, performance of the QA/QC samples, sampling surrogates, and internal standards.

Although an independent review of the data by the INEEL Sample Management Office (SMO) was not completed, the analytical data reports and data reduction calculations were reviewed by the contract laboratory Quality Officer, the Project Technical Leads, and the BBWI Project Quality Assurance Officer. All of the analytical data and offgas emissions results are judged to be useful for their intended purpose of completing an emissions inventory for the NWCF ETS system. The results are applicable to, and bounded by, the 2:1 volumetric blend of Tank WM-184 and Tank WM-181 feed composition, and NWCF ETS process operating parameters and conditions corresponding to the offgas sampling period.

#### Emissions Results

The concentration levels of the 20 highest VOC compounds emitted from the evaporator are plotted in Figure S-1. In general, volatile organic emissions are slightly higher at the start of an evaporator batch. The two highest volatile organics emitted from

the NWCF ETS were dodecane and acetone, which on a volumetric basis are only 50 ppbv and 30 ppbv, respectively. Acetone was also detected in the feed to the NWCF Calciner. Dodecane was not a target analyte for the liquid feed and therefore was not measured.

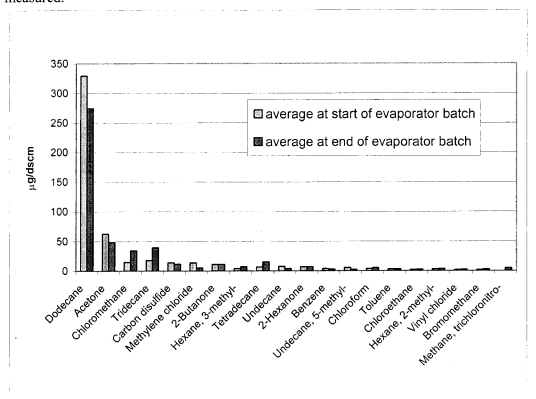


Figure S-1. Comparison of volatile organic emissions at the beginning and end of evaporator batches.

The top 20 SVOCs measured in the offgas stream are plotted in Figure S-2. SVOC emissions also appear to be slightly higher at the start of the evaporator batch. Benzoic acid (a target analyte) and benzaldehyde (a tentatively identified compound) were the two most prevalent semi-volatile organics emitted during operation of the NWCF ETS. The maximum emissions concentrations for benzoic acid and benzaldehyde were 310 ppbv and 80 ppbv, respectively.

Nearly all of the compounds are derivatives of benzene or other cyclic compounds and are possibly the products of incomplete combustion of the kerosene used to heat the Calciner. With the exception of benzoic acid, all of the SVOC species emitted from the evaporator were also detected during the NWCF Calciner offgas emissions inventory (Boardman 2001). It is therefore postulated that these compounds entered the Tank Farm System when Calciner scrub was recycled to the tank farm. They could also be formed by oxidation of benzene and toluene in the acidic waste solutions.

On a volumetric basis, the sum of all volatile and semi-volatile organics is less than 1 ppm. The hourly total emissions rate for all volatile and semi-volatile organic emissions was less than 0.02 lbs/hr.

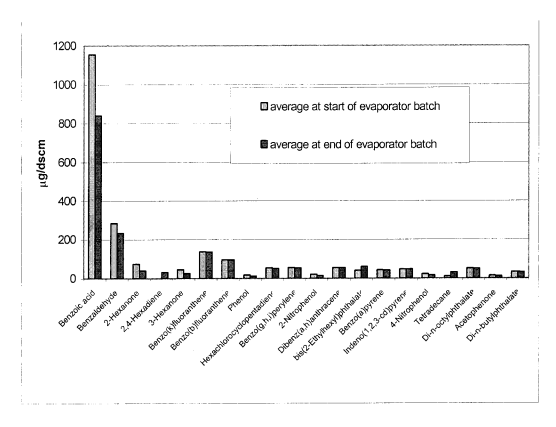


Figure S-2. Comparison of semi-volatile organic emissions at the beginning and end of evaporator batches.

The average metals emissions rates at the beginning and end of evaporator batches are plotted in Figure S-3. As anticipated, metals emissions, including mercury, were typically higher at the end of an evaporator batch when the evaporator solution reached its maximum density. The emissions of all metals species appear to correlate with the solution density.

Total particulate and chloride emissions rate averages at the start and end of two evaporator batches were very low. The sum of chloride emission contributions from HCl and Cl<sub>2</sub> was less than 1 ppmv. Particulate emissions were slightly higher at the beginning of the batch which followed the trend of the semi-volatile organic species emissions.

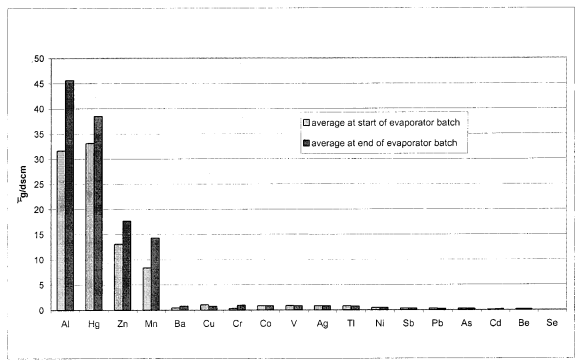


Figure S-3. Comparison of metals emissions at the beginning and end of evaporator batches (not including final probe rinse species apportionment).

#### **Emissions Risk**

The emission rate measurements were used to calculate the risk to human health. Pollutants from the NWCF ETS are released from the same point (*i.e.*, the INTEC main stack) and under the same conditions as NWCF Calciner emissions. Therefore, to a close approximation, the NWCF ETS hazards and risks can be scaled using the risk terms previously determined for the NWCF Calciner operations (Boardman 2001).

It was observed that the emissions rates, and hence component-specific risk contributions, were generally much lower from the NWCF ETS than from the NWCF Calciner. Most of the materials "found" were present at levels below the analytical laboratory reporting limits. The summed hazardous quotient (HQ) for all emissions from the NWCF ETS was 6.2e-6 as compared to the EPA target criteria of 0.25. The cancer risk was 1.3e-10 compared to an EPA target of 1e-5. The semi-volatiles were the largest contributor to the HQ and the Risk. The most significant species was a phthalate (bis(2 ethylhexyl)phthalate) which is a common contaminate from plastics present in laboratory and sampling areas.

In conclusion the measured emissions from the NWCF ETS are extremely low for all categories of pollutants. The estimated cancer risk and health hazard quotient are each several orders less than the limit normally allowed by EPA.

## **ACKNOWLEDGEMENTS**

This report is the result of a concerted effort on the part of several Company support organizations. The authors wish to acknowledge C. N. Woodall, Sr. Technical Specialist, for set up of the sample collection equipment and support during sample collection and shipping, and for chemical and waste management. R.M. Gifford is recognized for administrative support, training coordination, records management, and preparation of this report. The project manager for both the offgas and liquid sampling activities was L. J. Young, who supported technical planning, equipment and sample collection area setup, work authorization prerequisites, and ultimately, the sample collection activities.

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## **ACRONYMS**

ACS American Chemical Society

ALD Analytical Laboratory Department

APS atmospheric protection system

CAS Chemical Abstract Service

CEMS continuous emissions monitoring system

CFL central file location

CFR Code of Federal Regulations

COC chain of custody

CVAAS cold vapor atomic absorption spectroscopy

DF decontamination factor

D/F dioxins and furans

DCS Distributive control system

D.I. deionized

DOT Department of Transportation

DQOs data quality objectives

DWSD Drinking Water Standards Division

EDD electronic data deliverables

EMSL Environmental Monitoring Systems Laboratory

EPA U. S. Environmental Protection Agency

GC/MS gas chromatography/mass spectrometry

HCl hydrochloric acid

HEPA high-efficiency particulate air (filter)

HQ hazardous quotient

HRGC/HRMS high resolution gas chromatography/high resolution mass spectrometry

INEEL Idaho National Engineering and Environmental Laboratory

INTEC Idaho Nuclear Technology and Engineering Center

IS internal standard

L&V limitations and validation

LCS laboratory control samples

LCSD laboratory control samples duplicate

LET&D Liquid Effluent Treatment and Disposal

MCP Management Control Procedure (Company Document Indicator)

MDL method detection limit

MS/MDS matrix spike/matrix spike duplicate

NWCF New Waste Calcining Facility

PAH polycyclic aromatic hydrocarbons

PARCC precision, accuracy, representativeness, completeness, and comparability

PCBs polychlorinated biphenyls

PDS post digestion spikes

PEWE Process Equipment Waste Evaporator

PLN Plan (Company Document Designator)

PM particulate matter

PQAO Project Quality Assurance Officer

PTL Project Technical Lead(s)

QA/QC quality assurance/quality control

QAPjP quality assurance project plan

RAL Remote Analytical Laboratory

RCRA Resource Conservation and Recovery Act

RDL reliable detection limit

RFA/COC request-for-analysis/chain-of-custody

RL reporting limit (analytical laboratory established)

RPD relative percent difference

RPF relative potency factor

SAIC Science Applications International Corporation

SDG sample delivery group

SMO Sample Management Office

SMVOC sampling method for volatile organic compounds

SOW statement of work

STL Severn-Trent Laboratory- (Knoxville, Tennessee)

SVOC semi-volatile organic compound

SW Solid Waste

TAL target analyte list

TFF Tank Farm Facility

TICs tentatively identified compounds

TOC total organic carbon

TOS task order specific (statement of work)

TPR Technical Requirements Procedure (Company Document Designator)

VOC volatile organic compound

XX

# NWCF Evaporator Tank System 2001 Offgas Emissions Inventory

## 1. INTRODUCTION

Liquid wastes generated by fuel reprocessing and decontamination activities are stored in the Idaho Nuclear Technology and Engineering Center (INTEC) Tank Farm Facility. The Tank Farm wastes are currently being concentrated using the INTEC New Waste Calcining Facility (NWCF) Evaporator Tank System (ETS) (formally know as High Level Liquid Waste Evaporator HLLWE). The NWCF ETS currently operates under Resource Conservation and Recovery Act (RCRA) interim status. A RCRA Part B permit application for this unit is being prepared and is planned to be submitted in FY-2003. In order to support the permitting activities, the feed and output streams were characterized during evaporator operations in May and June, 2001. Characterization of the NWCF ETS process gaseous emissions were completed in conjunction with liquid feed and concentrated effluent analyses.

A detailed test plan (Test Plan for the HLLWE Effluent Gas Emissions Inventory, PLN-879) and quality assurance project plan (QAPjP- INTEC Quality Assurance Project Plan for the HLLWE Offgas Emissions Inventory Project, PLN-880) were developed for this project. The test plan discusses project organization, training requirements, safety implementation plans, sample collection objectives, and potential NWCF ETS offgas emissions. The QAPjP specifies the quality assurance and quality control (QA/QC) requirements, applicable quality standards, and both Idaho National Engineering and Environmental Laboratory (INEEL) and project-specific procedures for collecting, packaging, preserving, shipping, and analyzing the NWCF ETS offgas samples. The sample collection and analysis methods and procedures adhere to U.S. Environmental Protection Agency (EPA) protocol and technical requirements.

Science Applications International Corporation, Idaho Falls, Idaho (herein referred to as SAIC) was subcontracted to collect and recover the samples using the EPA prescribed procedures and equipment. SAIC also assisted BBWI in calculation of the air emissions rates using the data collected in the field and the sample analytical results. SAIC is recognized for its training and experience as a sample collection team. They previously supported the NWCF Calciner offgas emissions inventory project. Sample collection was performed using checklists and field data sheets.

Severn-Trent Laboratories, Knoxville, Tennessee (herein referred to as STL) performed the offgas sample preparations and analyses. The samples sent to STL were accompanied by a Request-for-Analysis Form (RFA), which documents the project-specific analytical specifications and quality control instructions to the laboratory. As part of the RFA documentation, a Chain-of-Custody (COC) and tractability record was maintained for all sample transfers to the laboratory. An analytical report for the final analytical data (STL 2001) was provided by STL. The analytical report includes a description of the analytical procedures that were used to acquire the data generated in support of this project.

Liquid feed streams and effluents associated with the NWCF ETS were collected in conjunction with the offgas sampling and were analyzed to complete mass balance and emissions inventory calculations. The samples were collected and analyzed under the Balance of Plant Sampling and Analysis Plan (inputs to Process Equipment Waste Evaporator (PEWE) and Liquid Effluent Treatment and Disposal (LET&D)). The liquid stream samples were collected and analyzed remotely to reduce operator and analyst exposure to radiation. Liquid sample collection was performed by the NWCF ETS operators

using double-needle sample collection system. The samples were sent the INTEC Remote Analytical Laboratory (RAL) for analyses.

The purpose of this report is to document and discuss the NWCF ETS offgas emissions inventory results and liquid feed stream analytical results. A technical description of the facility is followed by a description of the sample collection matrix and results. The risk associated with the offgas emissions has also been calculated and is presented herein.

## 2. NWCF ETS SYSTEM AND OFFGAS SAMPLING LOCATION

The NWCF ETS is a single-stage, thermal siphon, batch evaporator. Dilute Tank Farm liquid wastes are semi-continuously fed to the evaporator to maintain a constant level in the evaporator. A schematic of the NWCF ETS process is shown in Figure 1. The system consists of a feed tank (VES-NCC-152), a flash column (VES-NCC-150), a reboiler (HE-NCC-350), and a condenser (VES-NCC-151). Blended tank farm wastes are added to the flash column via the feed tank. When the level in the flash column reaches its normal operating level, steam is introduced into the shell side of the evaporator reboiler. As the evaporator solution temperature increases, its density decreases and the solution starts to rise. Steam bubbles form and further decrease the solution density. This draws the liquid from the bottom of the flash column into the tubes of the reboiler and creates a thermosiphon. The steam from the reboiler rises through a demister mesh and proceeds to the condenser. Typical NWCF ETS process operating conditions during sampling collection and analysis is shown in Table 1.

Once the desired specific gravity is reached, the concentrated solution is cooled and returned to the Tank Farm. The NWCF ETS overhead vapor is condensed in a total condenser and sent to the INTEC Process Equipment Waste Evaporator (PEWE) to be re-evaporated. Non condensable and purge gasses are vented from the feed tank and the condenser to the NWCF equipment vent system as shown in Figure 2. The equipment vent gasses join with the main process off-gas steam from the NWCF prior to the system high-efficiency particulate air (HEPA) filters.

Each batch generally requires 10-16 hours to process, followed by 10-16 hours to refill the feed tanks and to attend to waste transfers back to the Tanks Farm. Operating conditions that were monitored during emissions inventory testing are shown in discussed in Section 7. Normal operating conditions were maintained to provide the most stable and representative conditions throughout the sample collection period. All operating conditions are routinely recorded by the NWCF and Atmospheric Protection System (APS) control systems. These records are maintained by INTEC High Level Waste operations.

The offgas tie-in sample location used previously to sample the NWCF Calciner offgas stream was determined to be the best location for sample collection and offgas measurements for the scope and objectives of this project. The existing offgas tie-in location is downstream of the NWCF compressors and upstream of the APS. At this location, the offgas pipe is underground. The estimated offgas conditions at this location are listed in Table 1. Figure 3 shows the 12-inch ID pipe placement 9 ft underground, contained inside a larger 20 inch pipe encasement, which is inside a concrete encasement. The encasements provide the necessary physical protection and radiation shielding as the offgas flows to the APS.

Figure 4 shows a side view of the offgas tie-in location. This location is over 10 ft (10 pipe diameters) or more upstream and downstream of flow interference, so the flow should be reasonably straight (except for any disruption caused by the 12-inch tee). A 12-inch ID tee topped with a flange provides access through a manhole to the offgas pipe. Several penetrations (shown in top view in Figure 5) through the flange enable sample probe access and sample extraction.

The two-inch diameter port (line 2" POG-AR-156513) was used exclusively for the NWCF ETS offgas sample collection. This port is located at the centerline of the offgas duct cross section, allowing a vertical traverse of the duct to be made. A custom heated Method 5 probe (1.75 inch outside diameter) for was fabricated for sampling at this location. The sample probe was equipped with a compression fitting to provide a seal on the outer sheath of the sample probe. Pressurized air is used to continuously purge the annulus between the port inner wall and the probe sheath.

Table 1. Typical NWCF ETS operating conditions.

Table 1. Typical NWCF ETS operating of		X7.1 ()				
Parameter	DCS Identification Number	Value (a)				
HEPA filters						
Evaporator temperature	T150-1 through T150-10	95-110°C				
Steam to evaporator	F350-1C	1500-2000 lbs/hr				
Evaporator level	L150-1C	100-140 inches				
Evaporator density		1.0-1.35 g/mL				
Superheater (HE-NCC-335) outlet	T335-2C	150-205 °F				
offgas temperature (HEPA filter bank						
inlet temperature)						
HEPA filter inlet pressure	P130-2C	30 to 100 in. H <sub>2</sub> O				
HEPA filter stage 1 differential pressure	PD130-1-1C, -2-1C, -3-1C,	$0.5-10$ in. $H_2O$ (when online)				
-	-4-1C	0-0.5 in. H <sub>2</sub> O (when offline)				
Total differential pressure across HEPA	PD130-1C	2-18 in. H <sub>2</sub> O				
filter stages 1-3						
HEPA filter stage 3 outlet temperature	T130-1-1C, -2-1C, -3-1C,	80-150 °F				
	-4-1C					
NWCF process offgas flowrate (HEPA	F130-1C	50 - 1,000  scfm				
filter outlet offgas flowrate)						
	uipment Vent Conditions					
Offgas flow	F136-1C	500-1200 scfm				
Offgas temperature	T336-1C	60°- 80° F				
Offgas to APS pressure	P122-1	6-12 in. H <sub>2</sub> O vacuum				
Atmospheric Protection System (APS) an	d Other Equipment Downstream	of the Offgas Tie-in Sample				
Location	• •					
APS inlet offgas temperature (process	T-OGF-104	180-200°F				
offgas condenser outlet gas temperature						
APS inlet flowrate (process offgas flow)	F-OGF-2	1000-2000 scfm				
APS inlet offgas pressure	P-OGF-22	Negative 5-15 in. H <sub>2</sub> O (c)				
Main stack offgas flowrate	F-OGF-4/5	80,000-100,000 scfm				
Main stack offgas temperature	T-OGF-4-1, -5-1	70-100 °F				
a) If the value for an operating parameter dr	•	nge, or outside +/- 10% of the				

a) If the value for an operating parameter drifts outside of he indicated value range, or outside +/- 10% of the range is shown, then the test team leader must determine if sample collection should discontinue until NWCF operation is modified to correct the value.

b) Standard temperature and pressure is 60°F, 1 atmosphere.

c) This pressure is controlled using dampers on offgas blowers BLO-OGS-213 and -214, and can be adjusted to control the static pressure at the Offgas Tie-in location

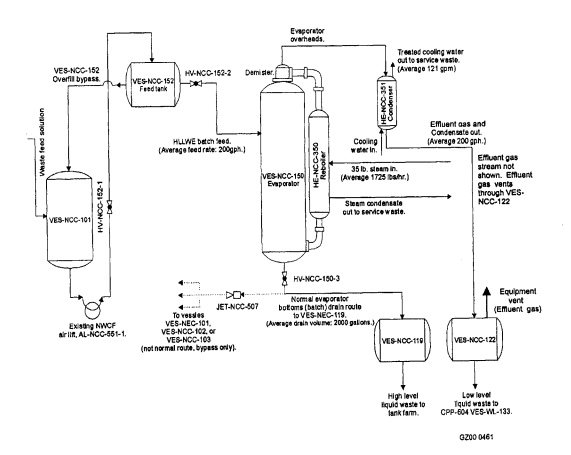


Figure 1. NWCF ETS system.

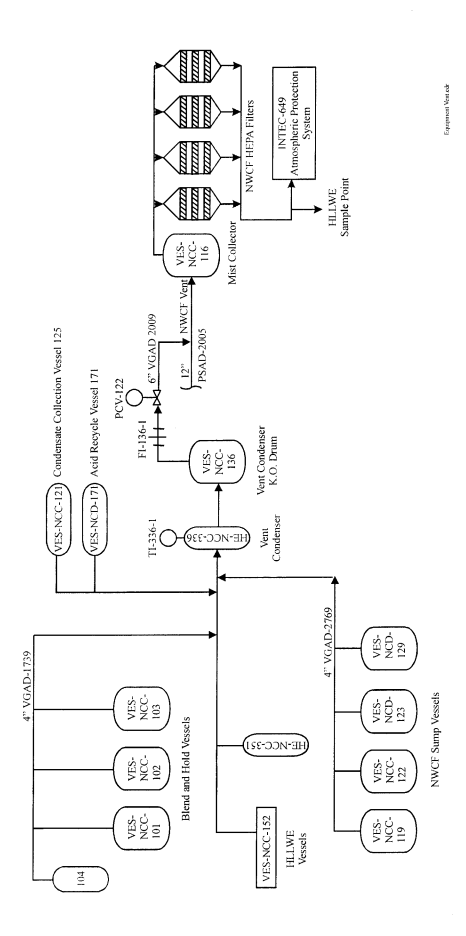


Figure 2. NWCF equipment vent.

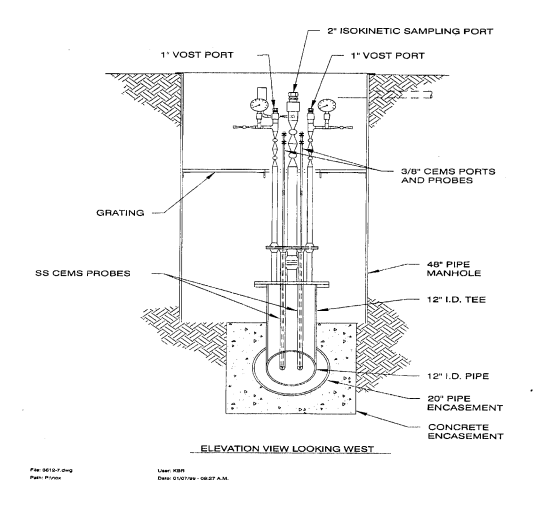
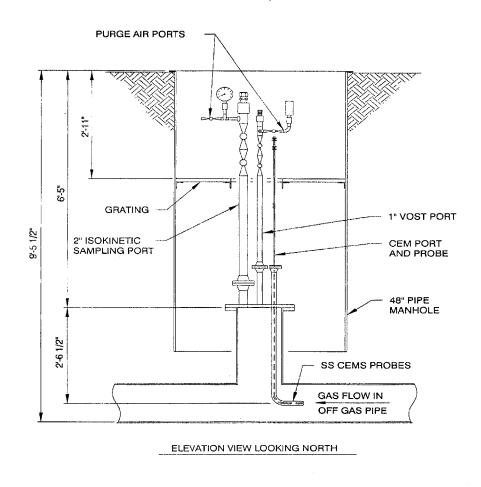


Figure 3. Offgas pipe axial view of the offgas tie-in sample location.

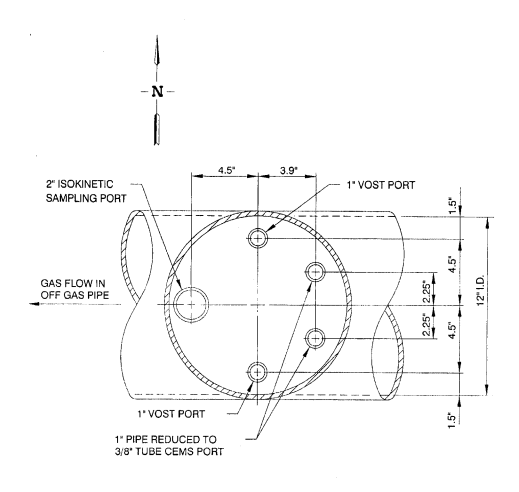


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Figure 4. Side view of offgas tie-in sample location.

This sample collection location does not meet all specified EPA requirements for offgas sample collection (40 Code of Federal Regulations (CFR) 60 Appendix A, Methods 1 and 2) since it is not possible to traverse the duct at two orthogonal positions. There may also be some mutual disturbance of the offgas flow pattern caused by the 3/8-inch tubes that are slightly upstream of the 2-inch access port as shown in a top view of the sample tie in (Figure 5). Fortunately, interference between sample ports is minimized as there is a clear path to oncoming gas flow as shown in the cross sectional view.

Another possible limitation to the sampling location is the presence of radionuclide contamination in the NWCF offgas duct. The procedure for inserting the 12 ft probe into the duct required donning of anti-contamination clothing and active monitoring by a Radiological Control Technician and Industrial Hygienist. The fragile probe tip can be easily damaged, and possibly could fall into the NWCF offgas duct- an event that is undesirable because it would introduction foreign material into the duct upstream of the Atmospheric Protect System. In addition, any potential presence of loose contamination in the duct could result in the spread of radiological contamination and possible exposure to the sample collection attendants. Hence, it was determined that the probe would be placed in the duct at a fixed point and not disturbed until the NWCF ETS offgas measurements were concluded. The probe was only articulated at the beginning of the sample collection tests in order to measure the vertical velocity profile in the duct.



### TOP VIEW OF SAMPLING TEE AT NWCF OFF-GAS TIE IN LOCATION

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Figure 5. Top view of offgas tie-in sample location.

## 3. SCOPE AND APPROACH

The purpose of this activity is to characterize the NWCF ETS process effluent gas emissions. The Tank Farm wastes are highly acidic (mainly nitric acid) inorganic salt solutions and contain several RCRA metals, including mercury, and trace amounts of volatile and semi-volatile hazardous organic compounds. Those components with low boiling points are released to the NWCF ETS offgas system during waste transfers, mixing, and evaporation. Trace amounts of heavy, nonvolatile hydrocarbons may also exist in the waste feed solutions; however, emissions of the nonvolatile hydrocarbons and also the nonvolatile metals is theorized to mainly occur by aerosol entrainment from the evaporator. The majority of the entrained droplets are believed to be collected in the offgas condensers, mist eliminators, and HEPA filters.

The scope of the NWCF ETS process effluent gas emissions inventory includes:

- Measurement of the NWCF offgas duct velocity, temperature, and flowrate during operation of the NWCF ETS
- Manual offgas sampling and analysis for particulate matter (PM), hydrochloric acid (HCl), Cl<sub>2</sub>, selected metals including Hg, volatile organic compounds (VOCs), and semi-volatile organic compounds (SVOCs)
- Measurement of radiological contaminate concentrations in sampling media using sample collection trains that are representative of the EPA sample collection trains
- Measurement of oxygen concentrations
- Measurement of probe rinseate for apportionment of metals and PM adsorbed on the probe to the respective train totals

Standard EPA sample collection and analysis methods were used to characterize the measure target analytes for each of the categories listed in Table 2. Measurements of moisture content and offgas temperature, velocity, and flowrate are included in each of the isokinetic sample train measurements. Sampling was conducted following EPA methodology with attention being given to the following:

- Development and adherence to a project quality assurance/quality control plan
- Implementation of sample chain-of-custody/requests-for-analysis, master sample lists, and sample labeling and tracking which assured in-field verifications of correctness of sample identifiers
- Development of target analyte lists (TAL) and precision, accuracy, representativeness, completeness, and comparability (PARCC) data quality indicators and objectives
- Collection of samples using checklists to record train set up, sample collection data, and sample recovery steps
- Collection and analysis of reagent blanks, trip blanks, and field blanks to assess prescribed data quality objectives
- Sample collection and documentation by a Project Quality Assurance Officer (PQAO)

- Application of EPA Solid Waste (SW)-846 and 40CFR 60 Appendix A reference methods for sample analysis
- Multiple reviews and verifications of field data, analytical data, process data, and resulting calculations of emissions rates

Table 2. Summary of NWCF ETS offgas sample collection and analysis methods.

Sample train procedure or Method	Measurement	Train description	Analytical procedures
2	Gas velocity, temp., swirl angle	S-type pitot, incline manometer, thermocouple	
0010	SVOCs	Isokinetic single-point, glass-lined probe, heated filter, XAD-2® sorbent, impingers	3542/8270C GC/MS (SVOCs) STL SOP KNOX-ID-0012
0060	Multiple metals including Hg	Isokinetic single-point, glass-lined probe, heated filter, impingers that contain sorbent solutions	6010A (ICAP) for multiple metals, 7470 (CVAAS) for mercury
0050 modified for PM	HCl, Cl <sub>2</sub> , PM		9056 / 9057 (IC for HCl, Cl <sub>2</sub> , and F); and Method 5 (gravimetric for PM), STL SOP KNOX-MS-0011
0031 SMVOC	VOCs	Single point, non-isokinetic, three sorbent tubes in series – (Tenax®/Tenax® /Anasorb® 747) and condensate trap	5041A/8260 GC/MS
3A or other (as requested by project lead)	$O_2$	Single point, nonisokinetic, heated sample line	Paramagnetism

The level of organics in the acidic Tank Farm waste solutions is very low and the NWCT ETS is operated at a much lower temperature than the NWCF Calciner. Therefore, because temperature and chemical precursors are not there in the system, it was determined that separate analysis of PCBs, and D/Fs was not necessary. It was determined that the results of the SVOCs for the target PAH compounds would be sufficient for risk assessment calculations. The offgas results presented in this report demonstrate that SVOC emissions are indeed negligible, as are precursors to PAHs and also higher

molecular weight compounds, including PAHs, PCBs, and D/Fs. Hence, the scope of the NWCF ETS offgas emissions inventory was limited to those methods shown in Table 2.

During the recent NWCF Calciner offgas emissions inventory (Boardman 2001), sample collection runs were conducted for analysis of 24 of the highly toxic semi-volatile polynuclear aromatic hydrocarbons (PAH), polychlorinated biphenyl compounds (PCB) and dioxins and furans (D/Fs), as well as SVOCs. This required separate runs with EPA Method 0010 and EPA Method 0023A trains. The samples are extracted and concentrated for subsequent analysis by high resolution gas chromatography/high resolution mass spectroscopy (HRGC/HRMS). Isotope dilution is used for each target analyte; thus, it was possible to achieve method detection limit concentrations for those analytes that were typically one-three orders of magnitude less than MDLS for the current reported project, where standard EPA Method 8270C gas chromatography/mass spectrometry (GC/MS) was specified. Lower detection limits for these compounds were desired to assist in the analysis of the Calciner performance, and also to provide the best possible data for Calciner emissions health risk assessment.

At the beginning of the sample collection period, the vertical gas velocity profile and swirl angle in the duct was measured to determine an appropriate fixed-point location to collect the offgas samples. Fixed-point sampling was necessary to avoid potential spread of contamination and possible damage to the glass probe tip. Two separate traverses were made to enhance accuracy of the velocity measurements. The minimum number of traverse points per Method 1 on the single (vertical) traverse for particulate and nonparticulate traverses (4) plus the pipe centerline were included in the traverses. The swirl angle at each traverse point and the average swirl angle per EPA Method 1, Section 2.4 were also determined. Subsequently, the probe was fixed at the point of maximum flow which corresponds EPA prescribed sample position at four inches from the pipe wall.

A set of two samples trains (referred to herein as "runs") were collected for each category of pollutants at the beginning and also at the end of evaporator runs. This provided a total of four runs for each method and brackets the emissions over the entire batch. One blank train (field blank) was also collected during the period that the four trains for each method were being run. Trip blanks and reagent blanks also were collected, as required by the QAPjP.

Oxygen concentrations were periodically recorded from the digital readout of the oxygen monitor located in the sampling tent during the manual sample collection operations. Because the off-gas sampled was essentially air supplied to ventilate NWCF vessels it was supposed and confirmed that the oxygen concentrations in the duct were similar to ambient air conditions. Therefore, it was determined that there was no need to constantly operate the oxygen monitor.

Moisture levels in the offgas were determined from the gravimetric and volumetric changes in the sample train resins and impingers, respectively. The moisture level was typically less than the dew point of the sample gas passing through the sample collection train condensers. At the most, only 1-2 mL of condensate was collected in any of the condensate knockout impingers. This is consistant with the use of a total condenser on the process and the addition of dry instrument air.

Finally, sample contamination survey trains were collected at the beginning of the tests in accordance with an INTEC management control procedure (MCP-1173, Package and Ship NWCF Offgas Emissions Samples Offsite for Analysis, Revision 2). Radioanlytical results from these trains were used to bracket the expected radioactivity range in the offgas samples to ensure that sample shipments to STL were in accordance with their radioactive materials license. Additional screening was performed routinely throughout the sample collection period. Every sample, as a minimum, was screened for gamma/beta emissions using a micro-R radiation detector, which is approximately 10 times more sensitive than the hand-held friskers used in the field by the Radiological Control Technicians. All of the

Method 5 filters were also submitted to the INTEC Radiochemical Laboratory for an extended duration (typically 12 hours) gamma count.

At the end of the sample collection period, the sample probe was removed from the duct and rinsed with acetone and nitric acid. These samples were surveyed for radiological contamination before being shipped to the analytical laboratory for measurement of PM and metals that were adsorbed on the probe. The results of these samples were used to address the technical acceptability of leaving the probe at a fixed position. It was assumed that the level of particulate in the duct would be negligible as a result of the sample location being downsteam of the NWCF HEPA filter banks. This was shown to be an appropriate assumption for the NWCF Calciner offgas emissions inventory project (Boardman 2001) and for the NWCF ETS as discussed later in this report. When a measurement of any target analyte in the probe rinsate was greater than the method detection limits, the result was apportioned to the metals and PM trains results.

Table 3 summarizes the sample collection trains, blank trains (field blanks), trip and reagent blanks, probe rinses, and radiological survey trains. Also listed is the sample collection date, time, and volume of offgas that was pulled through the train.

Table 3. Summary of samples collected in support of the INEEL NWCF ETS Effluent Gas.

Train ID or QC Sample No.	STL Sample No.'s Associated with this train	Target Analytes	Date Collected	Run Start Time	Run End Time	NWCF ETS Batch Num.	Reboiler Steam On Time	Reboiler Steam Off Time
SCS-EVAP-1	3269, 3270, 3271, 3272	Tritium, alpha/beta/gamma emitters	05/30/01	1030	1610	309	1026	1937
SCS-EVAP-2	3273, 3274, 3276, 3278	Tritium, alpha/beta/gamma emitters	05/31/01	0830	1240	310	0830	1747
0060-STRT-1	3279, 3280, 3281, 3282, 3283, 3284	Metals, including Hg	06/05/01	0930	1230	316	0951	1933
0060-STRT-2	3291, 3292, 3293, 3294, 3295, 3296,	Metals, including Hg	06/06/01	0751	1133	317	813	1715
0060 <sup>a</sup> Reagent Blanks	3297, 3298, 3299, 3300, 3301, 3348	Metals, including Hg	06/06/01	NA	NA	NA	NA	NA
0060-BT-1 <sup>b</sup> Blank Train	3302, 3303, 3304, 3305, 3306, 3307	Metals, including Hg	06/06/01	NA	NA	NA	NA	NA
0060-END-1	3326, 2227, 3328, 3329, 3330, 3331	Metals, including Hg	06/05/01	1600	1930	316	0951	1933
0060-END-2	3332, 3333, 3334, 3335, 3336, 3337	Metals, including Hg	06/06/01	1500	1802	317	0813	1715
Nitric Probe Rinse	334	Metals, including Hg	06/06/01	NA	NA	NA	NA	NA
0050-STRT-1	3308, 3309, 3310, 3311	PM, HCl, Cl <sub>2</sub> , HF, nitrate, nitrite	06/07/01	0800	1100	318	0800	1800

Table 3. Summary of samples collected in support of the INEEL NWCF ETS Effluent Gas.

Train ID or QC Sample No.	STL Sample No.'s Associated with this train	Target Analytes	Date Collected	Run Start Time	Run End Time	NWCF ETS Batch Num.	Reboiler Steam On Time	Reboiler Steam Off Time
0050-STRT-2	3312, 3313, 3314, 3315	PM, HCl, Cl <sub>2</sub> , HF, nitrate, nitrite	06/11/01	0750	1130	322	0821	1828
0050 <sup>a</sup> Reagent Blanks	3316, 3317, 3318, 3319, 3349	PM, HCl, Cl <sub>2</sub> , HF, nitrate, nitrite	06/07/01	NA	NA	NA	NA	NA
0050-BT-1 <sup>b</sup>	3322, 3323, 3324, 3325	PM, HCl, Cl <sub>2</sub> , HF, nitrate, nitrite	06/11/01	NA	NA	NA	NA	NA
0050-END-1	3338, 3339, 3340, 3341	PM, HCl, Cl <sub>2</sub> , HF, nitrate, nitrite	06/07/01	1405	1720	318	0800	1800
0050-END-2	3342, 3343, 3344, 3345	PM, HCl, Cl <sub>2</sub> , HF, nitrate, nitrite	06/11/01	1430	1730	322	0821	1828
Acetone probe rinse	3346	PM	06/11/01	NA	NA	NA	NA	NA
0010-STRT-1	3353, 3354, 3355, 3356, 3357, 3358	Semi-volatile organic compounds	06/18/01	0830	1130	329	0857	1819
0010-STRT-2	3372, 3373, 3374, 3375, 3376, 3377	Semi-volatile organic compounds	06/19/01	0830	1100	330	0824	1748
0010 <sup>a</sup> Reagent Blank	3378, 3343, 3444, 3445	Semi-volatile organic compounds	06/18/01	NA	NA	NA	NA	NA
0010-BT-1 <sup>b</sup>	3397, 3398, 3399, 3400, 3401, 3402	Semi-volatile organic compounds	06/18/01	NA	NA	NA	NA	NA

Table 3. Summary of samples collected in support of the INEEL NWCF ETS Effluent Gas.

Train ID or QC Sample No.	STL Sample No.'s Associated with this train	Target Analytes	Date Collected	Run Start Time	Run End Time	NWCF ETS Batch Num.	Reboiler Steam On Time	Reboiler Steam Off Time
0010-END-1	3403, 3404, 3405, 3406, 3407, 3408	Semi-volatile organic compounds	06/18/01	1500	1800	329	0857	1819
0010-END-2	3422, 3423, 3424, 3425, 3426, 3427	Semi-volatile organic compounds	6/19/01	1400	1715	330	0824	1748
0031-STRT-1	3359, 3360, 3361, 3362, 3363, 3364, 3365, 3366, 3367, 3368, 3369, 3370, 3371	Volatile organic compounds	06/20/01	0810	1048	331	0831	1748
0031-STRT-2	3379, 3380, 3381, 3382, 3383, 3384, 3385, 3386, 3387, 3388, 3389, 3390, 3391	Volatile organic compounds	06/21/01	0815	1152	332	0838	1749
0031 Field and Trip Blanks <sup>a,b</sup>	3392, 3393, 3394, 3395, 3396, 3441, 3442,	Volatile organic compounds	06/20/01	NA	NA	NA	NA	NA
0031-END-1	3409, 3410, 3411, 3412, 3413, 3414, 3415, 3416, 3417, 3418, 3419, 3420, 3421	Volatile organic compounds	06/20/01	1400	1715	331	0831	1748
0031-END-2	3428, 3429, 3430, 3431, 3432, 3433, 3434, 3435, 3436, 3437, 3438, 3439, 3440	Volatile organic compounds	06/21/01	1350	1710	332	0838	1749

a) Reagent blanks and trip blanks were obtained as identified in the master sample collection list.

b) Field QC samples are not exposed to the actual process offgas and thus are not correlated to ETS batches or run times. Probe rinses are not collected during active flow in the sampling slipstream, therefore, these are also not correlated to ETS batches or run times

## 4. RADIOLOGICAL SCREENING RESULTS

Two radiological contamination survey trains were collected at the beginning of the NWCF ETS offgas emissions inventory to establish the level of radiological contamination that could be uptaken by the EPA sample collection trains. The first train, identified as SCS-EVAP-1, was a hybrid of the Method 0060 for metals and Method 0050 for anions. The configuration of this train included a particle filter, followed by a condenser and condensate collection impinger, and then a pair of nitric acid/hydrogen peroxide impingers from the Method 0060 and a pair of sodium hydroxide impingers from the Method 0050 train. The acid and hydroxide impingers were used to capture the particulate and volatile radionuclides that are not disengaged by the filter and condensate trap. A gas volume of 3 dscm (dry, standard cubic meters) of gas was collected to match the volume of gas that was collected by the Method 0060 and 0050 sampling runs. Less than 2 mL of condensate was collected by this train- an insufficient amount for accurate analysis. Therefore, the condensate was added to the nitric/peroxide impinger solution.

The second radiological contamination train, identified as SCS-EVAP-2, was simply a standard Method 0010 train for semi-volatile organic collection, consisting of a particle filter, condenser, XAD-2® resin tube, a condensate trap and two organic-free water impingers. This train was mainly used to establish the level of contamination that could be potentially captured by the XAD-2® resin tube. The sample line, train glassware and filter housing were rinsed with acetone and methylene chloride. These rinses were composited into single sample for radiochemical analysis. The volume of gas collected was 3 dscm. The amount of condensate collected by the train was also very low (approximately 2 mL) for this train, indicating the offgas was essentially dry. The small amount of condensate was added to the organic-free impingers. Following the 12-hr gamma scan of the XAD-2® resin, the upper section of the resin bed, which first contract the sample gas and condensate, was extracted and prepped for gross alpha/gross beta counting.

Method 0031 for VOCs collection requires only 20 dsL (dry, standard liters) total, and only 5 dsL for each set of tubes. Therefore, the contamination levels established by the reference survey trains, at a total volume of 3000 dsL, clearly bounded the potential contamination picked up on the Tenax® resin tubes used in the Sampling Method for Volatile Organic Compounds (SMVOC) runs.

Analysis of the contamination survey train samples was completed by the INTEC Radiochemical Laboratory. Appropriate standards were prepared and used to provide quantitative results for the various sample collection media. Each sample was first analyzed by a non-intrusive gamma scan to measure gamma-emitting nuclides such as Ba<sup>137</sup> (which is the short lived daughter product of Cs<sup>137</sup>). Since Cs<sup>137</sup> is the most abundant non-volatile radionuclide in the waste, it is a convenient marker for the non-volatile radionuclides that could be present in the offgas samples, including Sr<sup>90</sup> and actinide isotopes. Therefore, an accurate gamma scan provides a basis for identifying the potential presence of Ba<sup>137</sup>, and hence Cs<sup>137</sup>, Sr<sup>90</sup>, and other fission products and actinides that may be present in the samples. A 12-hr gamma scan was performed to provide the most accurate analysis possible.

Following the gamma scan, the samples were prepared for gross alpha/gross beta counting. This required that the solid sample media be digested and then dried to obtain a valid measure of the particle emissions. The activity of tritium was determined by beta scintillation of an aliquot of the up-front liquid impinger contents to which the small amount of condensate was added. These fraction also absorb the largest percentage of the non-condensable water vapor.

Table 4 summarizes the radio-assay results for SCS-EVAP-1 and SCS-EVAP-1. Only an ultra low level of gross beta and gross alpha emissions was detected in the samples. The sample contamination

levels are conservatively less than the analytical laboratory sample screening acceptance criteria for Category I samples.

Table 4. Sample contamination survey train radio-assay results.

Train ID	Sample ID	Sample Media	Gamma	Gross Beta	Gross Alpha	Tritium
	3269	Particle filter	No nuclides identified	5.7E+00	9.8E-01	NA
				± 1.4E+00	± 6.7E-01	(dry sample)
		·		pCi	pCi	
	3270/3271	Condensate and nitric/peroxide	No nuclides identified	4.0E-02	Not detected	0.17
SCS-EVAP-1	composite	impinger contents		± 2.1E-02		μCi/sample
	;			pCi/mL		
	3272	Hydrogen peroxide impinger contents	No nuclides identified	4.3E+00	3.4E-02	NA (negligible
				± 2.1E-01	± 4.0E-02	condensate in sample)
				pCi/mL	pCi/mL	
	3273	Particle filter	No nuclides identified	1.85E+01	6.0E+00	NA
				± 4.0E+00	± 2.6E+00	(dry sample)
				pCi	pCi	:
	3276	XAD-2 <sup>®</sup> resin tube	No nuclides identified	5.84E00	Not detected	NA (negligible
				± 7.2E-01		condensate in sample)
				pCi/g		
SCS-EVAP-2	3278	Condensate and organic-free water	No nuclides identified	1.6E-02	Not detected	0.32
		impinger composite	Identified	± 1.9E-02		μCi/sample
				pCi/mL		
	3274	Organic solvent rinse composite	No nuclides identified	Not detected	Not detected	NA (negligible condensate in sample)

In order to ensure that the extremely low contamination levels in the offgas stream remained constant throughout the sample collection inventory, the particle filter for each Method 0060, Method 0050, and Method 0010 run, and one of the leading Tenax® tubes was submitted for a 12-hour gamma scan. This had little or no effect on sample preservation condition of the filters. Although the Tenax® was not maintained at the required temperature of 4°C, it is not likely that the VOC analysis results were

adversely affected since the tube was kept sealed during the gamma scan. The results of the on-going screening were consistent with the baseline results. No nuclides were identified by these extended gamma-scan analyses.

Each sample was also "smear-wiped" and counted to verify there was no detectable fugitive contamination on the surface of the sample containers. These additional screening efforts corroborated the results of the sample contamination survey trains and verified that each train did not collect any significant contamination throughout the 3-week sample collection period.

Finally, the volume of offgas sampled, as well as the amount of condensate collected by all of the sample runs, was checked and compared to the sample contamination survey trains. This ensured that the level of condensate, and hence the estimated amount of tritium in the respective samples, was consistent with the baseline results.

## 5. OFFGAS SAMPLE ANALYTICAL RESULTS

SAIC was responsible for setting-up, operating, and recovering the sample collection trains in the contamination containment hood. Once the samples were obtained, custody was transferred to the BBWI project principals for radiological screening and shipment to STL. The inorganic samples produced by each run were shipped to STL in Department of Transportation (DOT) approved fiber boxes with metal inner canisters. The organic samples were placed on ice and packaged in expanded, insulated coolers in order to maintain temperature preservation requirements. Chain-of-Custody and Requests-for-Analysis forms were used to track each sample. Shipments requiring preservative cooling were made using overnight delivery in order to ensure temperature preservation and analysis time limits were met. All of the samples meet the preservation and sample analysis time requirements without exception.

The results presented in this section are extracted from the Final Analytical Report provided by STL (STL 2001). Excerpts of the text and tables are included in this report to provide a single project summary document. Appendix A is a listing of the analytical lab certificates of analysis. These data were used to calculate the offgas emissions rates presented in Section 9.

STL tabulated train totals for each of the four EPA train runs that were made to characterize the NWCF ETS effluent gas emissions that are discharged through the NWCF offgas system. The results for each train component were summed to provide a run total for each target analyte. Although the laboratory data were reported down to the method detection limit (MDL), the project has implemented the reliable detection level (RDL) as the minimum value for risk calculations. The "RDL" is the detection level recommended by EPA. It is defined as 2.623 times the MDL (2.623 X MDL).

Significant figures for both the constituent fractions and the cumulative total were determined according to ASTM Standard E29-93a (1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications". Laboratory-assigned data qualifiers are displayed with each target analyte when required. The majority of these method-based flags are standardly defined flags among environmental laboratories. The data flags attached to the train totals represent the cumulative set of flags assigned to the result for each component that is included as part of the respective train totals. Data flags for individual component sample fractions were only carried through to the train totals when that particular train component result had an observable mathematical impact (based on significant figures as cited above) on the value of the "train totals" result for that compound.

When assigned, the "less than" (<) sign indicates that at least one sample fraction result included in the run total is either a "non-detect" value that has been evaluated down to the MDL of the measurement, or an estimated "hit" value that is below the RDL. In either case, the final analyte value for any fraction that has a laboratory result below the RDL is raised to the default RDL value, and the actual value for the respective analyte is judged to be less than conservative reported value. This same logic carries through to the summation of train fractions to arrive at train totals.

Additional project-specific train total flags are applied to the run total values that are not standard EPA data flags. These project-specific flags are specific to the NWCF ETS Offgas Emissions Inventory project and are defined as follows:

- An "N" flag indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
- A "P" flag indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.

• An "A" flag indicates that the compound was measured (detected) in all of the sampling train components, or fractions.

## 5.1 Volatile Organic Compounds

The standard U.S. EPA Method 0031 SMVOC sampling train configuration was used to collect samples of the NWCF ETS offgas for the assessment of volatile compounds. Each run used four sets of volatile organic adsorption resins tubes. Each set of tubes was comprised of two Tenax<sup>®</sup> tubes and one Anasorb 747<sup>®</sup> tube in series. The resin tubes were followed by a condensate trap that was used to capture the condensate captured by all four set of tube for a given run. The volume of offgas collected across each set of tubes was 5 L over a time period of approximately 30-40 minutes. Hence the total volume of gas for each run was approximately 20 L, collected over a time span of around 2.5 hours. The purpose of multiple tube sets was to integrate the sampling event over a period of time to better obtain representative data that characterizes the offgas emissions of the offgas stream sampled and is recommended by Method 0031. Each run produced 12 resin tubes and one condensate fraction that were stored on ice after they were removed from the train.

## 5.1.1 VOC Target Analyte List

The SMVOC samples were analyzed for the volatile organic compound target analytes given in Table 5. Analyses of SMVOC samples were completed using thermal desorption of the tubes onto a purge-and-trap device. SW-846 Method 5041B was implemented to carry out the thermal desorption. Method 8260B was implemented to analyze the desorbed analytes using GC/MS. The two SMVOC Tenax® tubes from a sample set were analyzed together, while the Anasorb 747® tube was separately analyzed.

A GC/MS library search was performed on each SMVOC sample (including the condensate samples) for non-target analytes, or tentatively identified compounds (TICs). The search was performed for the thirty (30) largest identifiable non-target compounds having a response that was at least 10% of the response of the nearest internal standard, which was spiked at 0.25  $\mu$ g. The library search was conducted against the National Bureau of Standards library of mass spectral data containing an estimated 75,000 compounds. The matching criteria includes a nominal 85% match of the mass spectral features, and analyst discretion of all identities reported.

Table 5. Volatile organic compound target analyte list.

VOC Target Analyte	CAS Registry Number
Acetone	67-64-1
Acrylonitrile	107-13-1
Benzene	71-43-2
Bromobenzene	108-86-1
Bromochloromethane	74-97-5
Bromodichloromethane	75-27-4
Bromoform	75-25-2
Bromomethane	74-83-9
2-Butanone	78-93-3
n-Butylbenzene	104-51-8
sec-Butylbenzene	135-98-8
tert-Butylbenzene	98-06-6
Carbon disulfide	75-15-0
Carbon tetrachloride	56-23-5
Chlorobenzene	108-90-7
Chlorodibromomethane	124-48-1
Chloroethane	75-00-3
Chloroform	. 67-66-3
Chloromethane	74-87-3
2-Chlorotoluene	95-49-8
4-Chlorotoluene	106-43-4
1,2-Dibromo-3-chloropropane	96-12-8
1,2-Dibromoethane	106-93-4
Dibromomethane	74-95-3
1,2-Dichlorobenzene	95-50-1
1,3-Dichlorobenzene	541-73-1
1,4-Dichlorobenzene	106-46-7
Dichlorodifluoromethane	75-71-8
1,1-Dichloroethane	75-34-3
1,2-Dichloroethane	107-06-2
1,1-Dichloroethene	75-35-4
cis-1,2-Dichloroethene	156-59-2
trans-1,2-Dichloroethene	156-60-5
1,2-Dichloropropane	78-87-5
1,3-Dichloropropane	142-28-9
2,2-Dichloropropane	594-20-7
1,1-Dichloropropene	563-58-6
cis-1,3-Dichloropropene	10061-01-5
trans-1,3-Dichloropropene	10061-02-6

Table 5. Volatile organic compound target analyte list.

VOC Target Analyte	CAS Registry Number
Ethylbenzene	100-41-4
Hexachlorobutadiene	87-68-3
2-Hexanone	591-78-6
Isopropylbenzene	98-82-8
p-Isopropyltoluene	99-87-6
Methylene chloride	75-09-2
4-Methyl-2-pentanone	108-10-1
Naphthalene	91-20-3
n-Propylbenzene	103-65-1
Styrene	100-42-5
1,1,1,2-Tetrachloroethane	630-20-6
1,1,2,2-Tetrachloroethane	79-34-5
Tetrachloroethene	127-18-4
Toluene	108-88-3
1,2,3-Trichlorobenzene	87-61-6
1,2,4-Trichlorobenzene	120-82-1
1,1,1-Trichloroethane	71-55-6
1,1,2-Trichloroethane	79-00-5
Trichloroethene	79-01-6
Trichlorofluoromethane	75-69-4
1,2,3-Trichloropropane	96-18-4
1,2,4-Trimethylbenzene	95-63-6
1,3,5-Trimethylbenzene	108-67-8
Vinyl chloride	75-01-4
m-Xylene & p-Xylene	136777-61-2
o-Xylene	95-47-6

### 5.1.2 VOC Analytical Results

Tabulated data summaries for the SMVOC data are given in Appendix A. These tables have been extracted from the STL Analytical Laboratory Final Report for this project. The run total (in total  $\mu$ g) for each analyte represents the sum of the amounts found in all of the SMVOC sets collected during each sampling run, including the amount of analyte found in the SMVOC condensate sample. The SMVOC condensate sample results were obtained by multiplying the observed concentration in mass/volume units ( $\mu$ g/L) by the final condensate volume (L) collected to obtain a result in units of mass ( $\mu$ g).

The Method 0031 SMVOC Tube Set Total (total  $\mu$ g/set) result consists of the sum of the analytical results for the two Tenax<sup>®</sup> resin tube contents (analyzed together) and the analytical result for the Anasorb 747<sup>®</sup> tube contents. The calculation is conducted as follows:

(Total  $\mu g$  on the Tenax  $^{\otimes}$  Tubes #1 and #2) + (Total  $\mu g$  on the Anasorb 747  $^{\otimes}$  Tube)

= Total  $\mu g$  on the Method 0031 SMVOC tube set

### 5.1.3 VOC Data Quality Assessment

The tubes were shipped to the analytical laboratory and analyzed within two week in accordance with EPA guidance and the QAPjP. The SMVOC samples were received at the laboratory in good condition. The samples were held on ice until the laboratory custodian checked the cooler temperatures and logged the samples at the laboratory.

All samples were processed through the analytical methods as planned, and analytical results were obtained for all of the expected analyses, with one exception. Low surrogate recoveries were obtained for sample A-3364 (The Anasorb 747® tube for Run 1, Set 2) and therefore the data for this sample are not usable. The results for all other samples meet the data quality objectives (DQOs) specified in the QAPjP and are therefore usable for the NWCF ETS offgas emissions inventory and risk assessment.

The toluene result for the Run 2, Set 2 Tenax®/Anasorb 747® tube appears to be an outlier. This particular result was over ten times higher than any other back-half result. Also, this result was over eight times higher than the corresponding front-half result. There were several other runs that exhibited higher toluene results for the back-half fraction than the front-half fraction. These results are not consistent with the results for other analytes, including benzene, which clearly did not break through the front-half SMVOC tubes. These inconsistent results imply that there was a source of fugitive contamination available to these tubes. Since the field and trip blanks did not generally exhibit toluene (only one front-half field blank and one back-half trip blank had "hits" for toluene), it appears that the sample tubes were exposed to environments containing some toluene that were not available to the field or trip blanks.

The methylene chloride results for Run 1, Set 1 and Set 2, were inconsistent with subsequent test runs. The methylene chloride result for Run 1, Set 1, was much higher than for any other analysis. The Run 1, Set 2 result was lower than the result for Run 1. The remaining results were lower in concentration, and appear to reside within three standard deviations of the mean value. Also, the acetone result for Run 1, Set 1, was the highest result for acetone that was found in any of the offgas samples. The cause of the high early results is assumed to be contamination of the sampling probe with acetone and methylene chloride. The SVOC train was used immediately prior to collection of the SMVOC samples. During this sampling event, the MM-5 probe (that was used for the SVOC train and had been rinsed with acetone and methylene chloride at the conclusion of SVOC sample collection) was used to collect the SMVOC samples. Since the same probe was rinsed with acetone and methylene chloride, it is likely that these solvents found in the offgas VOC samples originated in the equipment, not in the offgas.

The SMVOC runs results show a rapid decrease in methylene chloride in the second and third tube sets for the first run. This supports the supposition that the higher amounts of methylene chloride in the first run were an artifact of the field procedures. In spite of this supposition, the result was used when calculating the emissions health risk since the risk factor and emissions rate, although higher than actual emissions, does not impact the outcome of the cumulative emissions risk.

### 5.1.3.1 VOC Breakthrough Evaluation

The analysis scheme of the three-tube configuration of Method 0031 included individual analysis of each resin sample. The historical criterion for evaluating occurrences of SMVOC system breakthrough states that less than 30 percent by weight of an analyte should be detected on the back tube relative to the total amount observed on the front two tubes. That is, the Anasorb 747® resin tube should not contain more than 30 percent of the analyte total found on the front Tenax® resin samples. The criterion does not apply when less than 75 nanograms of an analyte are detected on the back trap. Additionally, the criterion does not apply when the analytes are the ultra-low boiling point analytes such as dichlorodifluoroethane, chloromethane, bromomethane, chloroethane, and vinyl chloride.

The SMVOC apparatus was operated under near optimum conditions during on-site sampling. The sample stream entered the first resin tube at a nominal 10°C as monitored by a thermocouple at the base of the condenser. A 20-liter sample was the maximum volume of gas pulled across the resin tubes, and the sampling rate was a standard SMVOC approach at approximately 0.50 liters per minute. Under these conditions the analytes were universally trapped on the Tenax® tubes except for the low boiling point analytes noted above. There are some anomalous results for acetone and toluene in which back-half fractions contained more than 30 percent of the front-half amount, and the total was more than 75 nanograms. These results appear to be derived from fugitive contamination sources, and do not represent breakthrough to the back half SMVOC tubes under these conditions. Acetone and toluene are solvents used during sampling for rinsing glassware and tubing. The reagent sources are assumed to be the source of the fugitive contamination.

### 5.1.3.2 VOC Blank Data Assessment

Several types of Method 0031 SMVOC blanks were evaluated during the offgas sampling analyses in order to assess the sampling and analytical environments for possible fugitive contamination sources. SMVOC field blanks were collected in order to assess the sampling train environments for possible fugitive contamination sources. Standard SMVOC trip blanks were also collected, as well as a deionized water trip blank. A comparison of the blank samples is shown in Table 6.

The SMVOC tube field blank results do not indicate field contamination by any target analytes except the three common laboratory contaminants, acetone, methylene chloride and toluene. The data for the SMVOC tube trip blanks also exhibit the presence of these three target analytes, in addition to bromomethane that was also observed on the Tenax® tubes, and dichlorodifluoromethane that was observed on the Anasorb 747® tube. The aqueous trip blank did not exhibit general contamination. The laboratory blanks associated with these samples also exhibit the presence of acetone and methylene chloride, but at levels too low to account for the observed levels of acetone in the trip blanks. Methylene chloride, acetone, and toluene are typically considered common laboratory contaminants during data validation. The laboratory method blank and field blank results do not, however, exhibit toluene.

Table 6. Comparison of SMVOC blank sample results.

Table 6. Comparison o	SMV		ank san	ipie re	SMV	OC.	SMV	OC.			T		Ι	
			SMV	OC					SMV	OC	SMV	OC	SMV	OC
	Tenax	( Pair			Tenax	Pair	Anason Field I			. m :				
	Field I	Blank	Anasor Field I		Field I	Blank	Tield I	Jiank	Tenax Pa		Anasor Trip E		D.I. Wat Bla	
	06/21	1/01ª	06/21	/01ª	06/21	/01ª	06/21	./01ª	06/20	)/01ª	06/20	/O1ª	06/22	2/01ª
	A-3	392	A-33	393	A-33	394	A-33	395	A-34	141	A-34	142	A-33	396
Analyte	(μ	g)	<b>(μ</b> ξ	g)	(μչ	g)	(μյ	g)	(με	g)	(μչ	g)	(μg/	/L)
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,												
Acetone	0.063	J,B	0.063	J,B	0.063	J,B	0.071	J,B	0.82	В	0.27	В	1.2	U
Acrylonitrile	0.29	U	0.29	U	0.29	U	0.29	U	0.29	U	0.29	U	12	U
Benzene	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.63	U
Bromobenzene	0.010	U	0.010	U	0.010	U	0.010	U	0.010	U	0.010	U	0.84	U
Bromochloromethane	0.015	U	0.015	U	0.015	U	0.015	U	0.015	U	0.015	U	0.66	U
Bromodichloromethane	0.011	U	0.011	U	0.011	U	0.011	U	0.011	U	0.011	U	0.79	U
Bromoform	0.019	U	0.019	U	0.019	U	0.019	U	0.019	U	0.019	U	0.60	U
Bromomethane	0.015	U	0.015	U	0.015	U	0.015	U	0.015	J	0.015	U	0.47	U
2-Butanone	0.10	U	0.10	U	0.10	U	0.10	U	0.10	U	0.10	U	2.2	U
n-Butylbenzene	0.016	Ú	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.60	U
sec-Butylbenzene	0.0084	U	0.0084	U	0.0084	U	0.0084	U	0.0084	U	0.0084	U	0.39	U
tert-Butylbenzene	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.34	U
Carbon disulfide	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.26	U
Carbon tetrachloride	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.47	U
Chlorobenzene	0.0084	U	0.0084	U	0.0084	U	0.0084	U	0.0084	U	0.0084	U	0.63	U
Chlorodibromomethane	0.015	U	0.015	U	0.015	U	0.015	U	0.015	U	0.015	U	0.68	U
Chloroethane	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.34	U
Chloroform	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.73	U
Chloromethane	0.013	U	0.013	U	0.013	U	0.013	U	0.013	U	0.022	J	0.26	U
2-Chlorotoluene	0.0047	U	0.0047	U	0.0047	U	0.0047	U	0.0047	U	0.0047	U	0.50	U
4-Chlorotoluene	0.0047	U	0.0047	U	0.0047	U	0.0047	U	0.0047	U	0.0047	U	0.50	U
1,2-Dibromo-3-chloropropane	0.029	U	0.029	U	0.029	U	0.029	U	0.029	U	0.029	U	1.0	U
1.2-Dibromoethane	0.020	U	0.020	U	0.020	U	0.020	U	0.020	U	0.020	U	1.0	U
Dibromomethane	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.71	U
1,2-Dichlorobenzene	0.020	U	0.020	U	0.020	U	0.020	U	0.020	U	0.020	U	0.45	U
1,3-Dichlorobenzene	0.010	U	0.010	U	0.010	U	0.010	U	0.010	U	0.010	U	0.52	U
1,4-Dichlorobenzene	0.014	U	0.014	U	0.014	U	0.014	U	0.014	U	0.014	U	0.55	U
Dichlorodifluoromethane	0.013	U	0.013	U	0.013	U	0.013	U	0.013	U	0.022	J	0.26	U
1.1-Dichloroethane	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.39	U
1,2-Dichloroethane	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.47	U
1,1-Dichloroethene	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.37	U
cis-1,2-Dichloroethene	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.50	U
trans-1,2-Dichloroethene	0.019	U	0.019	U	0.019	U	0.019	U	0.019	U	0.019	U	0.31	U
1,2-Dichloropropane	0.013	U	0.013	U	0.013	U	0.013	U	0.013	U	0.013	U	0.55	U
1,3-Dichloropropane	0.013	U	0.013	U	0.013	U	0.013	U	0.013	U	0.013	U	0.52	U
1,5-Dictioroproparie	0.019		0.017		10.017		0.019		L		L		0.52	

Table 6. Comparison of SMVOC blank sample results.

Table 6. Comparison o			ink sam	iple re										
	SMV	OC			SMV	OC	SMV	OC	C) (I)	00	CN 437	00	CMA	10C
	Tenax	Pair	SMV	OC	Tenax	Pair	Anasor	b 747	SMV	OC.	SMV	OC.	SMV	/UC
	Tonax		Anasor	b 747			Field E		Tenax Pa	ir Trip	Anasor	b 747	D.I. Wa	ter Trip
	Field E	Blank	Field E	Blank	Field I	Blank			Blar	ık	Trip B	lank	Bla	nk
	06/21	/01ª	06/21	/01ª	06/21	/01ª	06/21	/01ª	06/20	/01ª	06/20	/01ª	06/22	2/01ª
	A-33	92	A-33	93	A-33	394	A-33	95	A-34	41	A-34	42	A-33	396
Analyte	(μք	g)	(μք	<u>(</u> )	(με	g)	(μg	g)	(µg	;)	(μք	g)	(μg	/L)
2,2-Dichloropropane	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.26	U
1,1-Dichloropropene	0.020	U	0.020	U	0.020	U	0.020	U	0.020	U	0.020	U	0.37	U
cis-1,3-Dichloropropene	0.012	U	0.012	U	0.012	U	0.012	U	0.012	U	0.012	U	0.71	U
trans-1,3-Dichloropropene	0.015	U	0.015	U	0.015	U	0.015	U	0.015	U	0.015	U	0.66	U
Ethylbenzene	0.0092	U	0.0092	U	0.0092	U	0.0092	U	0.0092	U	0.0092	Ŭ	0.50	U
Hexachlorobutadiene	0.025	U	0.025	U	0.025	U	0.025	Ú	0.025	U	0.025	U	0.58	U
2-Hexanone	0.063	U	0.063	U	0.063	U	0.063	U	0.063	U	0.063	U	0.84	U
Isopropylbenzene	0.0063	U	0.0063	U	0.0063	U	0.0063	U	0.0063	U	0.0063	U	0.42	U
p-Isopropyltoluene	0.012	U	0.012	U	0.012	U	0.012	U	0.012	U	0.012	U	0.42	U
Methylene chloride	0.025	В	0.027	В	0.025	J,B	0.028	В	0.15		0.15		1.3	J,B
4-Methyl-2-pentanone	0.071	U	0.071	U	0.071	U	0.071	U	0.071	U	0.071	U	0.71	U
Naphthalene	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.26	U
n-Propylbenzene	0.0055	U	0.0055	U	0.0055	U	0.0055	U	0.0055	U	0.0055	U	0.52	U
Styrene	0.0068	U	0.0068	U	0.0068	U	0.0068	U	0.0068	U	0.0068	U	0.52	U
1,1,1,2-Tetrachloroethane	0.0097	U	0.0097	U	0.0097	U	0.0097	U	0.0097	U	0.0097	U	0.55	U
1,1,2,2-Tetrachloroethane	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.58	U
Tetrachloroethene	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.016	U	0.50	U
Toluene	0.0066	U	0.12		0.0066	U	0.0066	U	0.0066	U	0.066		0.66	U
1,2,3-Trichlorobenzene	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.26	U
1,2,4-Trichlorobenzene	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.58	U
1,1,1-Trichloroethane	0.022	U	0.022	U	0.022	U	0.022	U	0.022	U	0.022	U	0.42	U
1,1,2-Trichloroethane	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.52	U
Trichloroethene	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.017	U	0.47	U
Trichlorofluoromethane	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.018	U	0.26	U
1,2,3-Trichloropropane	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.025	U	0.84	U
1,2,4-Trimethylbenzene	0.0076	U	0.0076	U	0.0076	U	0.0076	U	0.0076	U	0.0076	U	1.0	U
1,3,5-Trimethylbenzene	0.0050	U	0.0050	U	0.0050	U	0.0050	U	0.0050	U	0.0050	U	0.45	U
Vinyl chloride	0.0066	U	0.0066	U	0.0066	U	0.0066	U	0.0066	U	0.0066	U	1.6	U
m-Xylene & p-Xylene	0.050	U	0.050	U	0.050	U	0.050	U	0.050	U	0.050	U	1.0	U
o-Xylene	0.0066	U	0.0066	U	0.0066	U	0.0066	U	0.0066	U	0.0066	U	0.60	U
a. This is the date of sample collection	on													

Acetone, methylene chloride, and toluene are used as rinsing solvents during Modified Method 5 sampling. These common solvents were present in the sampling area during the test series. Also, these compounds were observed in the field blanks at somewhat higher levels. Acetone was observed at a relatively high level in the SMVOC Tenax Pair Field Blank. These results indicate that fugitive contamination sources may have existed during the sampling and transport of these samples Per the SMVOC method restrictions, results from the SMVOC sampling were not blank corrected.

A practical approach was devised to use the front-half sample results for Run 1, Set 2 despite the fact that the corresponding back-half results are unusable. In order to use these results in the train total for the run, an estimate was made of the results for the corresponding back-half fraction. This estimate was obtained by averaging the results for the back halves of the remaining three sets of tubes for this run. This value is equivalent to calculating the back-half concentrations based on three (3) sets of tubes, or a total of 60 L of offgas. This approach has the benefit of not discarding valid data, in keeping with the general principle that all data obtained should be disclosed to the monitoring agencies.

## 5.1.3.3 VOC Internal Standard Recovery Assessment

Three internal standard compounds are spiked prior to thermal desportion of the SMVOC adsorbent tube. The same three internal standards were spiked into the aliquots of VOST condensate samples that were analyzed. These standards are used as a basis for the calculations of the concentrations of the target analytes and surrogates. A summary of the volatiles internal standard performance for all of the samples collected during Runs 1, 2, 3, and 4 are listed in Table 7. Method SW-8260 requires internal standard recovery to be at least 50 percent but not more than 200 percent of internal standard (IS) areas for the daily standard. Internal standard performance for all samples and all matrices was well within required limits (-50 to +100 percent difference relative to the IS areas for the daily standard) for all samples.

Table 7. Volatile Organic Compound (VOC) internal standard recoveries.  Percent Difference <sup>a</sup>						
			Per	cent Differ		
Field Sample No.	Run No.	Sample Description	Internal Standard #1 Fluorobenzene	Internal Standard #2 Chlorobenzene-d <sub>s</sub>	Internal Standard #3 1,4-Dichlorobenzene-d <sub>4</sub>	
A-3359/A-3360	0031-STRT-1	Tenax Tubes #1 & #2 (Set #1)	1.7	1.8	-4.6	
A-3362/A-3363	0031-STRT-1	Tenax Tubes #1 & #2 (Set #2)	3.2	-4.9	-14	
A-3365/A-3366	0031-STRT-1	Tenax Tubes #1 & #2 (Set #3)	-2.2	-10	-19	
A-3368/A-3369	0031-STRT-1	Tenax Tubes #1 & #2 (Set #4)	-6.1	-16	-23	
A-3379/A-3380	0031-STRT-2	Tenax Tubes #1 & #2 (Set #1)	-6.4	-12	-20	
A-3382/A-3383	0031-STRT-2	Tenax Tubes #1 & #2 (Set #2)	-8.2	-11	-19	
A-3385/A-3386	0031-STRT-2	Tenax Tubes #1 & #2 (Set #3)	-14	-23	-32	
A-3388/A-3389	0031-STRT-2	Tenax Tubes #1 & #2 (Set #4)	-15	-22	-32	
A-3409/A-3410	0031-END-1	Tenax Tubes #1 & #2 (Set #1)	-18	-23	-32	
A-3412/A-3413	0031-END-1	Tenax Tubes #1 & #2 (Set #2)	-15	-20	-30	
A-3415/A-3416	0031-END-1	Tenax Tubes #1 & #2 (Set #3)	-6.7	-8.1	-15	
A-3418/A-3419	0031-END-1	Tenax Tubes #1 & #2 (Set #4)	-1.8	-4.3	-9.1	
A-3428/A-3429	0031-END <b>-</b> 2	Tenax Tubes #1 & #2 (Set #1)	-25	-34	-37	
A-3431/A-3432	0031-END-2	Tenax Tubes #1 & #2 (Set #2)	<b>-2</b> 3	-32	-39	
A-3434/A-3435	0031-END-2	Tenax Tubes #1 & #2 (Set #3)	-26	-37	-44	
A-3437/A-3438	0031-END-2	Tenax Tubes #1 & #2 (Set #4)	-27	-38	-43	
A-3392	0031-STRT-2	Tenax Tube Pair Field Blank	-22	-30	-38	
A-3394	0031-STRT-2	Tenax Tube Pair Trip Blank	-18	-30	-39	
A-3441	0031-END-1	Tenax Tube Pair Field Blank	-17	-15	-28	
A-3361	0031-STRT-1	Anasorb 747 Tube (Set #1)	-16	-14	-24	
A-3364	0031-STRT-1	Anasorb 747 Tube (Set #2)	-28	-28	-33	
A-3367	0031-STRT-1	Anasorb 747 Tube (Set #3)	-28	-31	-36	
A-3370	0031-STRT-1	Anasorb 747 Tube (Set #4)	-27	-28	-28	
A-3381	0031-STRT-2	Anasorb 747 Tube (Set #1)	-36	-42	-46	
A-3384	0031-STRT-2	Anasorb 747 Tube (Set #2)	-25	-28	-29	
A-3387	0031-STRT-2	Anasorb 747 Tube (Set #3)	-24	-26	-28	
A-3390	0031-STRT-2	Anasorb 747 Tube (Set #4)	-17	-22	-25	
A-3411	0031-END-1	Anasorb 747 Tube (Set #1)	-13	-11	-17	
A-3414	0031-END-1	Anasorb 747 Tube (Set #2)	-14	-13	-21	
A-3417	0031-END-1	Anasorb 747 Tube (Set #3)	-16	-12	-21	
A-3420	0031-END-1	Anasorb 747 Tube (Set #4)	-19	-12	-20	

Table 7. Volatile Organic Compound (VOC) internal standard recoveries.

Table /. Volatile Of	rganic Compound	(VOC) internal standard reco	veries.				
				Perce	nt Diffe	rence	а
Field Sample No.	Run No.	Sample Description	Internal Standard #1	Fluorobenzene Internal Standard #2	Chlorobenzene-d <sub>5</sub>	Internal Standard #3	1,4-Dichlorobenzene-d4
A-3430	0031-END-2	Anasorb 747 Tube (Set #1)	-16	,	-25	-	.33
A-3433	0031-END-2	Anasorb 747 Tube (Set #2)	-24		-29	-	35
A-3436	0031-END-2	Anasorb 747 Tube (Set #3)	-20	)	-30	-	∙38
A-3439	0031-END-2	Anasorb 747 Tube (Set #4)	-24		-39	-	33
A-3393	0031 <b>-</b> STRT-2	Anasorb 747 Field Blank	-20		-28	-	36
A-3395	0031-STRT-2	Anasorb 747 Trip Blank	-21		-25	-	31
A-3442	0031-END-1	Anasorb 747 Field Blank	-19		-22	_	32
A-3371	0031-STRT-1	VOST Condensate	-9.3	3	-7.0	_	13
A-3391	0031-STRT-2	VOST Condensate	-10		-8.8	_	15
A-3421	0031-END-1	VOST Condensate	-7.2	2	-5.0		9.8
A-3440	0031-END-2	VOST Condensate	-8.1		-5.6	_	11
A-3396	0031-STRT-2	VOST D.I. Water Trip Blank	-12		-9.3	-	15

a. Recoveries of internal standard compounds are not typically calculated for samples analyzed by Method 8260B and 8270C. Percent Difference is calculated using the following equation.

$$Percent \ Difference (\%D) = \frac{Observed \ Value - Expected \ Value}{Expected \ Value} x 100\%$$

Where: Observed Value = the area of the internal standard in the sample and

Expected Value = the area of the internal standards in the daily standard

## 5.1.3.4 VOC Surrogate Recovery Assessment

Four surrogate compounds were spiked onto all of the VOST samples before the thermal desorption process was initiated. The surrogate recoveries for the NWCF ETS offgas samples are presented in Table 8. Surrogate recoveries are within the targeted acceptance range (percent recovery between 50-150%), meeting the project DQOs except for sample A-3364. Low surrogate recoveries were obtained for sample A-3364 (The Anasorb 747® tube for Run 1, Set 2) and therefore the data for this sample are not usable. The results for Sample A-3364 indicate normal recoveries for the internal standard compounds, but very low recoveries for the surrogate compounds.

Table 8. Volatile Organic Compound (VOC) surrogate compound recoveries.

				Percent	Recoverya	
Field Sample No.	Run No.	Sample Description	Dibromofluoromethane	1,2-Dichtoroethane-d <sup>b</sup>	Toluene-d <sub>s</sub>	Bromofluorobenzene
A-3359/A-3360	0031-STRT-1	Tenax Tubes #1 & #2 (Set #1)	84	81	101	83
A-3362/A-3363	0031-STRT-1	Tenax Tubes #1 & #2 (Set #2)	90	85	114	99
A-3365/A-3366	0031-STRT-1	Tenax Tubes #1 & #2 (Set #3)	88	84	113	98
A-3368/A-3369	0031-STRT-1	Tenax Tubes #1 & #2 (Set #4)	85	80	115	92
A-3379/A-3380	0031-STRT-2	Tenax Tubes #1 & #2 (Set #1)	88	83	111	95
A-3382/A-3383	0031-STRT-2	Tenax Tubes #1 & #2 (Set #2)	91	86	110	97
A-3385/A-3386	0031-STRT-2	Tenax Tubes #1 & #2 (Set #3)	81	75	108	85
A-3388/A-3389	0031-STRT-2	Tenax Tubes #1 & #2 (Set #4)	87	80	114	90
A-3409/A-3410	0031-END-1	Tenax Tubes #1 & #2 (Set #1)	84	79	109	85
A-3412/A-3413	0031-END-1	Tenax Tubes #1 & #2 (Set #2)	85	80	109	91
A-3415/A-3416	0031-END-1	Tenax Tubes #1 & #2 (Set #3)	83	78	101	82
A-3418/A-3419	0031-END-1	Tenax Tubes #1 & #2 (Set #4)	87	86	103	80
A-3428/A-3429	0031-END-2	Tenax Tubes #1 & #2 (Set #1)	87	79	116	87
A-3431/A-3432	0031-END-2	Tenax Tubes #1 & #2 (Set #2)	79	74	105	86
A-3434/A-3435	0031-END-2	Tenax Tubes #1 & #2 (Set #3)	83	77	117	90
A-3437/A-3438	0031-END-2	Tenax Tubes #1 & #2 (Set #4)	90	82	121	92
A-3392	0031-STRT-2	Tenax Tube Pair Field Blank	76	70	102	86
A-3394	0031-STRT-2	Tenax Tube Pair Trip Blank	58	49 <sup>1</sup>	60	41 <sup>1</sup>
A-3441	0031-END-1	Tenax Tube Pair Field Blank	82	77	95	86
A-3361	0031-STRT-1	Anasorb 747 Tube (Set #1)	83	79	94	77
A-3364	0031-STRT-1	Anasorb 747 Tube (Set #2)	7.9 <sup>1</sup>	6.4 <sup>1</sup>	2.91	3.6 <sup>1</sup>
A-3367	0031-STRT-1	Anasorb 747 Tube (Set #3)	106	100	121	92
A-3370	0031-STRT-1	Anasorb 747 Tube (Set #4)	86	83	99	82
A-3381	0031-STRT-2	Anasorb 747 Tube (Set #1)	96	88	117	86
A-3384	0031-STRT-2	Anasorb 747 Tube (Set #2)	73	71	90	74

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Table 8. Volatile Organic Compound (VOC) surrogate compound recoveries.

		npound (VOC) surrogate comp	Percent Recovery <sup>a</sup>			
Field Sample No.	Run No.	Sample Description	Dibromofluoromethane	1,2-Dichloroethane-d <sup>b</sup>	Toluene-d <sub>s</sub>	Bromofluorobenzene
A-3387	0031-STRT-2	Anasorb 747 Tube (Set #3)	77	73	92	74
A-3390	0031-STRT-2	Anasorb 747 Tube (Set #4)	73	68	96	79
A-3411	0031-END-1	Anasorb 747 Tube (Set #1)	87	84	101	84
A-3414	0031-END-1	Anasorb 747 Tube (Set #2)	91	86	107	86
A-3417	0031-END-1	Anasorb 747 Tube (Set #3)	91	88	104	88
A-3420	0031-END-1	Anasorb 747 Tube (Set #4)	78	75	96	81
A-3430	0031-END-2	Anasorb 747 Tube (Set #1)	83	76	105	88
A-3433	0031-END-2	Anasorb 747 Tube (Set #2)	88	79	110	82
A-3436	0031-END-2	Anasorb 747 Tube (Set #3)	82	75	108	85
A-3439	0031-END-2	Anasorb 747 Tube (Set #4)	85	79	110	87
A-3393	0031-STRT-2	Anasorb 747 Field Blank	77	68	99	73
A-3395	0031-STRT-2	Anasorb 747 Trip Blank	82	74	92	75
A-3442	0031-END-1	Anasorb 747 Field Blank	83	77	98	81
A-3371	0031-STRT-1	VOST Condensate	102	104	106	106
A-3391	0031-STRT-2	VOST Condensate	100	103	107	106
A-3421	0031-END-1	VOST Condensate	101	101	106	105
A-3440	0031-END-2	VOST Condensate	100	103	106	105
A-3396	0031-STRT-2	VOST D.I. Water Trip Blank	102	107	107	106
Laboratory 1	 Target Recovery Ra	nge for Tenax <sup>®</sup> & Anasorb 747 <sup>®</sup> Tubes:	50-150	50-150	50-150	50-150
***************************************	Laboratory Targe	et Recovery Range for Aqueous Samples:	80-120	80-120	80-120	72-135

a. Percent Recovery is calculated using the following equation:

Percent Re cov ery (%R) = 
$$\frac{Observed\ Value}{Expected\ Value} \times 100\%$$

Where: Observed Value = the measured mass of the surrogate standard in the sample and Expected Value = the mass of the surrogate standard spiked into the sample.

This percent recovery is outside of the laboratory target recovery range.

### 5.1.3.5 VOC Analytical Data Quality Assessment

The sampling and analytical objectives expected for this data set were to present an acceptable characterization of the project target volatile organic compounds from the NWCF ETS offgas. The data quality indicators collectively indicate that the sampling and analytical processes for the SMVOC samples were in control during the sampling runs. Data have been collected and reviewed that allow the relative precision and accuracy to be measured for the target analytes. The data quality indicators indicate that most of the data are of acceptable quality, and that sufficient data has been obtained to characterize the project target volatile organic compounds from the NWCF ETS offgas.

There were several indications that fugitive emissions may have been present during sampling. Acetone, methylene chloride and toluene were present in at least some of the field and trip blanks. Some inconsistency of the methylene chloride results has been discussed. Toluene exhibited results that simply do not make sense, particularly with respect to higher levels of toluene that were detected in some of the back half samples. Results for these particular three constituents are not considered to demonstrate that these constituents are truly in the ETS offgas at these concentrations. These are the major classical environmental laboratory sample preparation solvents that are often detected as contaminants in sample results.

The only serious quality control deficiency was the low surrogate recovery for the Tenax<sup>®</sup>/Anasorb 747<sup>®</sup> sample for Run 1 (Set 2). This deficiency is adequately handled by the substitution of average backhalf results from the other Run 1 tube sets.

## 5.2 Semi-volatile Organic Compounds

A standard U.S. EPA Method 0010 (Modified Method 5, or MM-5) sampling train configuration was used to collect samples of the NWCF ETS offgas for the assessment of semi-volatile organic compounds (SVOCs). A total nominal volume of 3.0 dscm of offgas was sampled in each run over 3-4 hours. The Method 0010 SVOC train configuration is comprised of six fractions:

- particulate filter
- solvent rinse of the front half of the filter holder, the sampling probe and the nozzle
- XAD-2<sup>®</sup> resin tube,
- solvent rinse of the back half of the filter holder, the coil condenser and connecting glassware,
- composite sample containing the stack gas condensate and impinger contents, and
- impinger and connecting glassware solvent rinses.

A trip/reagent blank was collected and a set of blank train (field blank) samples were analyzed to assess extraneous sources of contamination available to these samples.

# 5.2.1 SVOC Target Analyte List

Analyses of SVOC samples were completed per SW-846 Methods 3542 and 8270C by first extracting the samples with methylene chloride, then analyzing the extracts using GC/MS. The SVOC target analytes are listed in Table 9.

Table 9. SVOC target analyte list.

Analyte	CAS Registry Number
Acenaphthene	83-32-9
Acenaphthylene	208-96-8
Acetophenone	98-86-2
Aniline	62-53-3
Anthracene	120-12-7
Benzidine	92-87-5
Benzoic acid	65-85-0
Benzo(a)anthracene	56-55-3
Benzo(a)pyrene	50-32-8
Benzo(b)fluoranthene	205-99-2
Benzo(ghi)perylene	191-24-2
Benzo(k)fluoranthene	207-08-9
Benzyl alcohol	100-51-6
bis(2-Chloroethoxy)methane	111-91-1
bis(2-Chloroethyl)ether	111-44-4
bis(2-Ethylhexyl)phthalate	117-81-7
4-Bromophenyl-phenylether	101-55-3
Butylbenzylphthalate	85-68-7
Carbazole	86-74-8
4-Chloro-3-methylphenol	59-50-7
4-Chloroaniline	106-47-8
2-Chloronaphthalene	91-58-7
2-Chlorophenol	95-57-8
4-Chlorophenyl phenyl ether	7005-72-36
Chrysene	218-01-9
Di-n-butylphthalate	84-74-2
Di-n-octylphthalate	117-84-0
Dibenz(a,h)anthracene	53-70-3
Dibenzofuran	132-64-9
1,2-Dichlorobenzene	95-50-1
1,3-Dichlorobenzene	541-73-1
1,4-Dichlorobenzene	106-46-7
3,3'-Dichlorobenzidine	91-94-1
2,4-Dichlorophenol	120-83-2
Diethylphthalate	84-66-2
Dimethyl phthalate	131-11-3
2,4-Dimethylphenol	105-67-9

Table 9. SVOC target analyte list.

Analyte	CAS Registry Number
4,6-Dinitro-2-methylphenol	534-52-1
2,4-Dinitrophenol	51-28-5
2,4-Dinitrotoluene	121-14-2
2,6-Dinitrotoluene	606-20-2
I,2-Diphenylhydrazine	122-66-7
Fluoranthene	206-44-0
Fluorene	86-73-7
Hexachlorocyclopentadiene	77-47-4
Hexachlorobenzene	118-74-1
Hexachlorobutadiene	87-68-3
Hexachloroethane	67-72-1
Indeno(1,2,3-cd)pyrene	193-39-5
Isophorone	78-59-1
2-Methylnaphthalene	91-57-6
2-Methylphenol	95-48-7
3-Methylphenol & 4-Methylphenol	65794-96-9
N-Nitroso-di-n-propylamine	621-64-7
N-Nitrosodimethylamine	62-75-9
N-Nitrosodiphenylamine	· 86-30-6
Naphthalene	91-20-3
2-Nitroaniline	88-74-4
3-Nitroaniline	99-09-2
4-Nitroaniline	100-01-6
Nitrobenzene	98-95-1
2-Nitrophenol	88-75-5
4-Nitrophenol	100-02-7
2,2'-Oxybis(1-chloropropane)	108-60-1
Pentachlorobenzene	608-93-5
Pentachloronitrobenzene	82-68-8
Pentachlorophenol	87-86-5
Phenanthrene	85-01-8
Phenol	108-95-2
Pyrene	129-00-0
Pyridine	110-86-1
1,2,4-Trichlorobenzene	120-82-1
2,4,5-Trichlorophenol	95-95-4
2,4,6-Trichlorophenol	88-06-2
1,2,4,5-Tetrachlorobenzene	95-94-3

The samples were also analyzed for non-target organic compounds as directed in the QAPjP. A GC/MS library search was performed on each SVOC sample (including the condensate samples) for TICs (tentatively identified compounds). The search was performed for the thirty largest non-target compounds that exhibited a response greater than 10% of the response of the nearest internal standard (the extract is spiked at 20  $\mu$ g/mL). The standard extract volume was 1.0 mL; hence, the TICs were reported down to a level of 2  $\mu$ g when the original extract was not diluted (dilution factor or DF = 1). The backhalf composite sample extracts for Runs 1 through 4 were analyzed at a five-fold dilution (DF = 5); therefore, TICs in these fractions were only reported down to 10  $\mu$ g.

The library search was conducted against the National Bureau of Standards library of mass spectral data containing an estimated 75,000 compounds. The matching criteria included a nominal 85% match of the mass spectral features, and analyst discretion of all identities reported. TICs that were derived from column bleed, surrogate addition, or aldol condensation were excluded from the report. Also, compounds that were reported as SMVOC target compounds were not reported as semi-volatile TICs because the SMVOC method provides more reliable data for these compounds.

### 5.2.2 SVOC Analytical Results

The particulate filter was combined with its associated solvent rinses to form a "front-half" composite sample. The XAD-2 resin tube was combined with its associated solvent rinses to form a "back-half" composite sample. The stack gas condensate, impinger contents and associated glassware rinses were also combined to form a composite sample. These three fractions are analyzed separately. Unique data quality control indicators are used for each fraction.

Tabulated data summaries that present the SVOC data are given in Appendix A. These tables have been extracted from the STL Analytical Laboratory Final Report for this project. For each Method 0010 offgas sampling run, the "SVOC Run Total" for each analyte (in mass units of  $\mu g$ ) represents the sum of the amounts found in all of the SVOC fractions collected during that run.

### 5.2.3 SVOC Data Quality Assessment

The sample fractions were sent to STL via overnight express mail to ensure that sample preservation and analysis schedules required by the QAPjP would be met. All of the samples were received by the laboratory in good condition. Sample extractions were performed within the requirements specified in the QAPjP.

On the basis of all the quality assurance indicators, all of the semi-volatile organic compound data obtained from the SVOC runs are usable and representative of the NWCF ETS offgas contents. The only deficiencies in accuracy and precision indicated by the laboratory control samples (LCS) and matrix spike samples appear to be unrelated to sample data quality.

Blank sample data indicate that sources of fugitive contamination available to the NWCF ETS offgas samples were minimized. Only phthalate esters were found in significant concentrations in the blanks. Although the back half composite containing the XAD-2<sup>®</sup> sample extracts were somewhat hostile to the internal standard compounds, the analysis of these extracts at two levels of dilution appears to provide a reliable assessment of the offgas contents.

Recoveries of the surrogates indicate that the preparation and analysis processes during the SVOC sample determinations were in control with respect to all of the analytes for the offgas sample analyses. The surrogate recoveries are within the prescribed acceptance ranges and do not indicate any bias to the data. Sample dilution was required to achieve acceptable recovery of three of the six internal standards,

and this had the affect of increasing detection limits for those analytes that are correlated to the recovery of these standards.

### 5.2.3.1 SVOC Blank Data Assessment

A standard SVOC trip/reagent blank (unused, sealed XAD-2® resin tube) and a blank train run samples was collected to assess potential fugitive contamination sources in the sampling environment. Review of the SVOC blank indicates that very little contamination due to fugitive emissions exists in the samples as a result of storage or transport of the sample collection media. The trip blank data exhibited low levels of acetophenone and 1,4-dichlorobenzene that are below the standard laboratory reporting limit (RL). Several tentatively identified compounds were identified in the trip blank data. Notably, benzaldehyde, methyl benzaete, and ethyl benzaldehyde were found, along with several miscellaneous hydrocarbons. These compounds were probably artifacts of the XAD-2® medium or the transport and storage of the samples, and were not found in the laboratory method blanks.

Review of the SVOC blank train results indicates that little contamination of the samples occurred as a result of sample handling or contact with the MM-5 sampling train components. Acetophenone and 1,4-dichlorbenzene were found at low levels that were similar to the trip blank, and may have originated in the sampling media, or were possibly introduced to the media during transport and storage of the samples. The target analytes found in the blank train samples also included di-n-butylphthalate, bis-2-ethylhexylphthalate, and di-n-octylphthalate. The phthalate esters are considered common laboratory contaminants, and are commonly found in certain plastics and plastic tubing. These compounds were not found in the laboratory method blanks, so their origin appears to be with the sampling process. The TICs found in the back-half composite sample of the blank train were similar in identity and concentration to those found in the trip/reagent blank. Benzaldehyde, methyl benzoate, ethyl benzaldehyde, and several miscellaneous hydrocarbons were observed at levels that were similar to the trip/reagent blank results. The origin of these contaminants may have been either the sampling media or the transport and storage of the samples. The front-half and impinger composite samples exhibited low concentrations of some additional TICs that are not found in the back-half composite samples.

### 5.2.3.2 SVOC Internal Standard Recovery Assessment

Internal standards are used as the basis for calculation of the concentrations of the target analytes and surrogates. Six IS compounds were spiked into all of the sample extracts prior to analysis. Method SW-8270 required limits, in terms of percent difference relative to the IS area for the lab's daily standard, are -50% to +100%. The internal standard responses for the front-half composite sample extracts, and the condensate and impinger contents composite sample extracts were acceptable and do not indicate any deficiency in data quality. Also, the quality assurance samples exhibit acceptable recoveries of the internal standards. The only cases of significantly reduced recovery of the internal standards are noted for the back-half fractions of the offgas samples which include the XAD-2® resin. A summary of the semi-volatile internal standard performance is given in Table 10.

Table 10. SVOC train sample internal standard compound recoveries.

14010 101 01 00	l am sample	internal standard compound recove		Percent Difference (%) <sup>a</sup>				6) <sup>a</sup>	
Field Sample No.	NWCF ETS Sample ID	Sample Description	Analytical Dilution Factor	Internal Standard 1 1,4-Dichlorobenzene-d <sub>4</sub>	Internal Standard 2 Naphthalene-d <sub>8</sub>	Internal Standard 3 Acenaphthene-d <sub>10</sub>	Internal Standard 4 Phenanthreene-d <sub>10</sub>	Internal Standard 5 Chrysene-d <sub>12</sub>	Perylene-d <sub>12</sub>
A-3353/A-3354	0010-STRT-1	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	1	9.2	7.5	-5.6	-16	-27	-22
A-3355/A-3356	0010-STRT-1	XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser	5	-28	-29	-61 <sup>b</sup>	-38	-57	-100 <sup>b</sup>
		Solvent Rinses	100	7.0	3.2	-1.6	.0.62	3.5	-50
A-3357/A-3358	0010-STRT-1	Condensate, Impinger Contents, and Glassware Solvent Rinses	1	13	12	0.16	-4.4	-20	-16
A-3372/A-3373	0010-STRT-2	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	1	12	6.9	-1.1	-7.0	-24	-20
A-3374/A-3375	0010-STRT-2	XAD-2 Resin Tube/Back Half of the	5	2.4	-5.2	-25	-14	-27	-100 <sup>b</sup>
		Filter Holder and Coil Condenser Solvent Rinses	100	2.4	1.9	-1.2	-3.2	-1.0	-24
A-3376/A-3377	0010-STRT-2	Condensate, Impinger Contents, and Glassware Solvent Rinses	l	-15	-18	-16	-16	-26	-24
A-3397/A-3398	0010-STRT-2	Blank Train Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	1	5.3	2.4	-5.8	-11	-24	-22
A-3399/A-3400	0010-STRT-2	Blank Train XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser Solvent Rinses	1	-0.69	-7.8	-18	-21	-34	-27
A-3401/A-3402	0010-STRT-2	Blank Train Condensate, Impinger Contents, and Glassware Solvent Rinses	1	7.2	9.3	-1.9	-9.8	-22	-18
A-3403/A-3404	0010-END-1	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	1	7.6	9.6	0.92	-13	-26	-20
A-3405/A-3406	0010-END-1	XAD-2 Resin Tube/Back Half of the	5	-6.6	-5.7	-37	-14	-30	-100 <sup>b</sup>
		Filter Holder and Coil Condenser Solvent Rinses	100	1.7	1.8	1.4	-2.4	3.1	-65 <sup>b</sup>
A-3407/A-3408	0010-END-1	Condensate, Impinger Contents, and Glassware Solvent Rinses	1	0.38	4.2	-2.7	-8.3	-22	-20
A-3422/A-3423	0010-END-2	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	1	5.7	4.1	-2.2	-13	-28	-29
A-3424/A-3425	0010-END-2	XAD-2 Resin Tube/Back Half of the		14	9.9	-28	0.05	-19	-100 <sup>b</sup>
		Filter Holder and Coil Condenser Solvent Rinses	100	-2.9	-2.0	-6.5	-6.0	-3.7	-66 <sup>b</sup>
A-3426/A-3427	0010-END-2	Condensate, Impinger Contents, and Glassware Solvent Rinses	1	1.8	4.2	-3.1	-8.7	-20	-16
A-3378	0010-STRT-2	XAD-2 Resin Tube Trip/Reagent Blank	1	-14	-15	-22	-20	-29	-27
	-	Internal Standard Laboratory Percent Difference Acceptance Limits:		-50 to 100	-50 to 100	-50 to 100	-50 to 100	-50 to 100	-50 to 100

Table 10.	SVOC tra	ain sample i	nternal standa	ard compound	l recoveries.

					Per	cent Diff	erence (	(%) <sup>a</sup>	
Field Sample No.	NWCF ETS Sample ID	Sample Description	Analytical Dilution Factor	Internal Standard 1 1,4-Dichlorobenzene-d <sub>4</sub>	Internal Standard 2 Naphthalene-d <sub>8</sub>	Internal Standard 3 Acenaphthene-d <sub>10</sub>	Internal Standard 4 Phenanthreene-d <sub>10</sub>	Internal Standard 5 Chrysene-d <sub>12</sub>	Perylene-d <sub>12</sub>

a. Recoveries of internal standards are not typically calculated for samples analyzed by Method 8260B and 8270C, and internal standard recoveries are evaluated as Percent Difference from the daily standard. Percent Difference is calculated using the following equation:

Percent Difference (%D) = 
$$\frac{Observed\ Value - Expected\ Value}{Expected\ Value} \times 100\%$$

Where: Observed Value = the area of the internal standard in the sample and

Expected Value = the area of the internal standard in the daily standard.

b. This value is outside of the laboratory and project target acceptance range.

The sample extracts of the back-half fractions of the offgas sample trains were analyzed at a dilution factor of five since analysis of the extracts without dilution gave evidence of loss of several of the internal standards. The internal standard perylene-d12 did not recover from the extract analyses at a dilution factor of five. There was insufficient recovery of this internal standard to allow quantification of the related target analytes without applying further dilution. The internal standard compounds acenaphthene-d10 and chrysene-d12 also exhibited recoveries that are lower than the target acceptance criteria in the 1:5 analysis of the extract for the back-half fraction extract for Run 1. However, there was sufficient recovery of each of these internal standards to provide useful results for the related target analytes.

A second extract analysis was performed for each of the offgas back-half composite samples at a dilution factor of 100. The back-half composite sample extracts exhibit reduced recovery of perylene-d12 at the increased dilution factor of 100, but there was sufficient recovery of perylene-d12 to quantify the seven target analytes that are calculated relative to it. The results based on perylene-d12 for the 1:100 dilution of the extracts are usable.

### 5.2.3.3 SVOC Surrogate Recovery Assessment

Six surrogate compounds were spiked onto all of the SVOC samples before extraction per SW-846 Method 8270C. Three of the surrogates are base/neutral compounds, and three of the surrogates are acid extractable. All of the three acid extractable surrogates are phenols, which are a class of organic compounds that contain a benzene ring with the hydroxyl group attached. A sampling surrogate compound was also applied to the XAD-2® resin tubes at the laboratory before sampling. This additional surrogate provides a measurement of the efficiency of the entire process, from sampling to the conclusion of the analysis. The sampling surrogate applied to the XAD-2® tubes used to collect samples was <sup>13</sup>C<sub>3</sub>-naphthalene. This is a base-neutral compound that is distinguished from the native naphthalene by carbon-13 labeling. A summary of the semi-volatile surrogate performance is given in Table 11.

Table 11. SVOC surrogate compound recoveries.

			Percent Recovery (%) <sup>a</sup>						
Field Sample No.	NWCF ETS Sample ID	Sample Description	2-Fluorophenol	Phenol-d <sub>5</sub>	Nitrobenzene-d <sub>5</sub>	2-Fluorobiphenyl	2,4,6- Tribromophenol	Terphenyl-d <sub>14</sub>	Surrogate
A-3353/A-3354	0010-STRT-1	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	71%	66%	66%	70%	75%	76%	NA
A-3355/A-3356	0010-STRT-1	XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser Solvent Rinses	65%	50%	87%	148% <sup>b</sup>	44%	122%	78%
A-3357/A-3358	0010-STRT-1	Condensate, Impinger Contents, and Glassware Solvent Rinses	73%	75%	80%	86%	84%	93%	NA
A-3372/A-3373	0010-STRT-2	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	65%	61%	64%	66%	70%	76%	NA
A-3374/A-3375	0010-STRT-2	XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser Solvent Rinses	68%	55%	84%	106%	52%	96%	86%
A-3376/A-3377	0010-STRT-2	Condensate, Impinger Contents, and Glassware Solvent Rinses	39%	35%	43%	46%	53%	82%	NA
A-3397/A-3398	0010-STRT-2	Blank Train Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	75%	68%	73%	76%	74%	81%	NA
A-3399/A-3400	0010-STRT-2	Blank Train XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser Solvent Rinses	69%	66%	74%	81%	3.1% <sup>b</sup>	82%	78%
A-3401/A-3402	0010-STRT-2	Blank Train Condensate, Impinger Contents, and Glassware Solvent Rinses	67%	71%	72%	79%	75%	89%	NA
A-3403/A-3404	0010-END-1	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	72%	67%	69%	71%	72%	78%	NA
A-3405/A-3406	0010-END-1	XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser Solvent Rinses	79%	57%	90%	128% <sup>b</sup>	50%	100%	86%
A-3407/A-3408	0010-END-1	Condensate, Impinger Contents, and Glassware Solvent Rinses	77%	80%	77%	78%	81%	90%	NA
A-3422/A-3423	0010-END-2	Particulate Filter/Front Half of the Filter Holder and Probe Solvent Rinses	75%	71%	74%	76%	66%	81%	NA
A-3424/A-3425	0010-END-2	XAD-2 Resin Tube/Back Half of the Filter Holder and Coil Condenser Solvent Rinses	68%	55%	92%	134%	54%	110%	85%
A-3426/A-3427	0010-END-2	Condensate, Impinger Contents, and Glassware Solvent Rinses	67%	76%	78%	81%	83%	88%	NA
A-3378	0010-STRT-2	XAD-2 Resin Tube Trip/Reagent Blank	55%	61%	63%	67%	0.0% <sup>b</sup>	84%	66%
		Surrogate Laboratory Percent Recovery Acceptance Range:	%001-61	15-124%	35-122%	34-115%	33-130%	28-132%	50-150%

a. Percent recovery of surrogate compounds is calculated using the following equation:

Percent Re cov ery (%) =  $\frac{Observed \ Value}{x \ 100!}$ 

Expected Value

where: Observed Value = the measured mass of the surrogate standard in the sample and

Expected Value = the mass of the surrogate standard spiked on the sample.

b. This value is outside of the laboratory and project target acceptance range.

The laboratory surrogate recoveries that were obtained for the front-half composite samples are generally excellent. There are no obvious problems with the analysis of this matrix, which includes the particulate filter and solvent rinses of the front-half of the filter holder and the probe. All of the laboratory surrogate percent recovery values for the front-half samples were all within the target acceptance range. There were no significant differences between the acid surrogate performance for Runs 1 through 4 and the blank train.

The back-half samples exhibited good recoveries for both the base-neutral and the acid extractable surrogates. 2-Fluorobiphenyl recovery from the back-half fractions of Runs 1, 3 and 4 were above the target acceptance range for surrogate recovery, while the percent recovery of 2-fluorobiphenyl for Run 2 was near the upper end of the target acceptance range. The percent recovery of 2-fluorobiphenyl for both the blank train and the XAD-2<sup>®</sup> resin tube trip/reagent blank was near the center of the target acceptance range. Reduced recovery of the related internal standard, acenapthylene-d<sub>10</sub> in the offgas samples appears to cause the observed high recoveries of 2-fluorobiphenyl. A high bias to the results for all of the target analytes that are calculated against acenaphthylene-d<sub>10</sub> appears to be indicated by the increased recovery of 2-fluorobiphenyl. Inspection of the data shows that there are no positive results based on acenaphthylene-d<sub>10</sub>. Therefore, the detection limits are defensible and there is no adverse impact on data quality.

Phenol-d<sub>5</sub> and 2-fluorophenol exhibited acceptable recovery in the blank train back-half sample and the XAD-2 trip/reagent blank, but 2,4,6-tribromophenol did not recover well. The 2,4,6-tribromophenol results were much lower than the target acceptance limits, with no recovery at all from the trip/reagent blank sample. These results were atypical of the data set in that the 2,4,6-tribromophenol recoveries in the offgas samples were well within the target acceptance range for all four runs. The associated laboratory method blank exhibited a good recovery of the 2,4,6-tribromophenol, but the laboratory control sample (LCS) and the laboratory control sample duplicate (LCSD) each exhibited a low recovery of this surrogate. Also, the LCS have low recoveries of 4-nitrophenol and pentachlorophenol, which are acidic compounds potentially sensitive to pH affects. There was no control of pH for these solid matrix samples. The surrogates were applied to the matrix in a Soxhlet extractor immediately prior to extraction. There were low levels of nitrates present in the offgas, and it is probable that the XAD-2 samples were rendered slightly acidic by contact with the offgas. Acidic samples are more likely to release the acid extractable surrogates during the extraction process. Recovery of the 2,4,6-tribromophenol was well within the target acceptance limits for the offgas samples; hence, there is no impact on the offgas data quality because the phenomenon is not observed in the offgas sample results.

On the basis of laboratory surrogate recovery, the results for the SVOC train back-half composite sample data were usable for assessment of the NWCF ETS offgas contents. All of the laboratory surrogate recoveries are consistently within the target acceptance range. The condensate, impinger contents and associated glassware rinse samples also exhibited acceptable results for all of the laboratory surrogate compounds. The sampling surrogate compound that was applied to the XAD-2® resin tubes at the laboratory before sampling provide an independent measurement of the efficiency of the entire process, from sampling to the conclusion of the analysis. The  $^{13}C_3$ -naphthalene sampling surrogate recovered well from all runs. This further indicates that the data for the SVOCs are reliable.

## 5.2.3.4 SVOC Laboratory Control Sample and Matrix Spike Sample Performance

Laboratory control samples associated with the front half (glass fiber filter) matrix were prepared and analyzed in duplicate as required by the QAPjP. There are two spiked compounds that exhibited results that were outside the target acceptance limits. The recovery of 2,4-dinitrotoluene in the LCS was slightly below the target acceptance range. The relative percent difference (RPD) for pentachlorophenol was above the target acceptance range. However, neither of these results indicates a significant loss of

data quality. All of the laboratory surrogates in the LCS/laboratory control samples duplicate (LCSD) analyses associated with the front-half samples are within target acceptance ranges.

Laboratory control samples associated with the back-half composite samples were prepared and analyzed in duplicate. The base-neutral surrogate recovery results for the LCS were similar to the results for the field samples, and were within the stated target acceptance range. The LCS exhibit low recoveries of 4-nitrophenol and pentachlorophenol, which are acidic compounds that are sensitive to pH effects. Pentachlorophenol is the most acidic of the spiked compounds, and had almost no recovery. The laboratory control sample and laboratory control sample duplicate both have low recoveries of the acidic laboratory surrogate 2,4,6-tribromophenol. This appears to be because there was no control of pH for these solid samples. The surrogate recoveries of similar compounds were well within target acceptance ranges for the NWCF ETS offgas samples; consequently, there is no impact on the offgas data quality as a result of these low recoveries in the LCS/LCSD.

Laboratory control samples based upon the aqueous matrix were prepared and analyzed in duplicate. The condensate and impinger contents sample for Run 2 was split into three aliquots, and a matrix spike/matrix spike duplicate pair was analyzed. All spiked analyte and surrogate percent recovery and RPD results for the laboratory control and matrix spike samples were within the stated target acceptance ranges.

### 5.3 Metals

A standard U.S. EPA Method 0060 Multi-Metals Train (MMT) configuration was used to collect NWCF ETS offgas samples for the assessment of metals (including mercury) content. An offgas sample having a nominal volume of 3.0 m³ was collected over a duration of 3-4 hours. Two sample collection runs were completed at the beginning and two at the end of consecutive evaporator batches to provide a total of four runs to characterize metals emissions. The following sample components were collected from the 0060 train after the completion of each run:

- Particle Filter
- 0.1N Nitric Acid (HNO<sub>3</sub>) Probe Rinse
- 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> Impinger Contents
- Empty Impinger 4 Contents
- 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> Impingers
- 8N HCl Impinger Rinses

Final nitric acid probe rinse samples were collected after the sampling was completed. The glass-lined sampling probe could not be routinely removed from the offgas sampling manifold for rinsing after each run. Instead, the front-half rinse samples were limited to the connecting tubing, filter housing elements and other connecting glassware that were installed outside the manifold assembly. The glass-lined probe was only removed from the offgas line after all offgas trains were completed for the test series. Probe rinse samples of the glass-lined probe were collected to assess the maximum amount of metals that adhered to the inside of the probe during testing. The probes were rinsed three times with acetone, followed by three rinses with 0.1N nitric acid. These samples were combined for metals analysis after the acetone probe rinses were analyzed for PM. Since the same probe was used for all sampling runs used during the test series, a larger volume of offgas was represented by these acetone and nitric acid probe rinses than was pulled through the probe just for metals analysis.

Field reagent and blank samples were collected in compliance with the QAPjP. Laboratory method blanks were also prepared and analyzed in support of the data, as required by SW-846 sample analysis requirements.

### 5.3.1 Metals Target Analyte List

The target analyte list for the metals is given in Table 12. These metals, except for Hg, were analyzed by inductively coupled argon plasma spectroscopy per EPA Method 6010B. Mercury (Hg) was analyzed using cold vapor atomic absorption spectroscopy (CVAAS) Method 7470A.

Table 12. Metals target analyte list.

Aluminum (Al)	Chromium (Cr)	Nickel (Ni)
Antimony (Sb)	Cobalt (Co)	Selenium (Se)
Arsenic (As)	Copper (Cu)	Silver (Ag)
Barium (Ba)	Lead (Pb)	Thallium (Tl)
Beryllium (Be)	Manganese (Mn)	Vanadium (V)
Cadmium (Cd)	Mercury (Hg)	Zinc (Zn)

### 5.3.2 Metals Analytical Results

The compatible matrices of the MMT samples were separated into the train's front-half and backhalf samples during the sample preparation steps. The front-half samples consisted of the 0.1N nitric acid probe and filter housing rinses, and the particulate filter. The combined offgas condensate, and the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> impinger catches comprised the MMT back-half composite. The train's fourth impinger was used to separate the components of the train that contained 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> from the peroxide contained in the front impingers. The fourth impinger was left empty during set-up. Its contents at the completion of a run and 0.1N HNO<sub>3</sub> collection rinses were only analyzed for mercury. The 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> solution is used to trap elemental mercury (Hg). Thus, a composite sample was prepared from these impingers for the analysis of mercury, only. The internal surfaces of the backhalf impinger glassware received a final deionized (D.I.) water rinse during the train sample collections. These rinsates were collected and added to the appropriate impinger sample and included in the back-half composites. A final rinse of the potassium permanganate impingers was conducted with 8N HCl, and was collected as a separate sample for mercury analysis.

The sample collection and analysis are outlined in the STL final report. Sample fraction and run totals are listed in Appendix A. The run totals (in  $\mu g$ ) are the sum of results for the front half and backhalf composite samples, and in the case of mercury, include the permanganate/sulfuric impinger composite totals. Results for the blank trains were not used to correct train totals as allowed by Method 0060 guidance.

### 5.3.3 Metals Data Quality Assessment

Samples were shipped using overnight delivery service to the laboratory. All Method 0060 train samples that were collected during the offgas sampling program were received at the laboratory in good condition. No losses of samples due to breakage or loss of shipment occurred. Samples derived from the Method 0060 trains are not required to be refrigerated after sample collection. All of the run fractions were processed and analyzed by STL in the time requirements specified in the QAPjP.

The data quality indicators of the sampling and analytical processes for the Method 0060 train samples indicate that the metals data collected from these samples represent acceptable characterization of the offgas emissions. Sufficient data has been collected to allow accuracy and precision to be measured for these parameters. Accuracy has been measured by analyzing LCS, post digestion spikes (PDS), and a limited set of matrix spikes. Precision was measured by the preparation and analysis of laboratory control sample duplicates. The data quality indicators present sufficient evidence that the data are of acceptable quality and are usable for the NWCF ETS offgas emissions inventory.

### 5.3.3.1 Metals Trains Reagent and Blank Data Assessment

Laboratory method blanks were prepared and analyzed in support of the data. Review of the laboratory method blank data indicates that the laboratory did not appear to introduce significant fugitive contamination to the samples.

Reagent blanks were collected in the field and processed to assess the inherent metallic analyte background in the media being used for sampling. A representative front-half composite reagent blank (a quartz fiber particulate filter and 120 mL 0.1 N HNO<sub>3</sub> rinse reagent), a representative back-half reagent blank (175 mL of the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> impinger solution), and a sample of the INTEC-supplied deionized water (210 mL) were collected and analyzed for the project target metallic analytes including mercury. Portions of the 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> impinger solution, and the 8N HCl rinse solution were collected and analyzed for mercury. Aluminum and selenium were detected in the front-half composite reagent blank at levels that were above the laboratory RL. The amounts of aluminum, antimony, barium, nickel, selenium, and zinc indicate that the reagent may have contributed to the offgas sampling train front-half results. Other metals were detected at relatively insignificant amounts. The back-half reagent blank exhibits manganese and zinc above the laboratory RL. The levels of aluminum, barium, chromium, nickel, lead, manganese, selenium, and zinc indicate that the reagent may have contributed to the offgas sampling train back-half fraction results. The INTEC deionized water did not exhibit significant levels of any metals, although aluminum was found at 5.2 μg.

The blank train samples exhibit similar results as those obtained for the reagent blanks, with two exceptions. Manganese was found in the blank train back-half composite sample at a level (3700  $\mu$ g) that far exceeded the offgas sample results. This occurrence can be traced to contamination of the back-half composite sample by KMnO<sub>4</sub> reagent. Zinc was found at a level of 34  $\mu$ g. This amount is roughly an order of magnitude higher that the reagent blank sample, and similar to the offgas sampling train results.

### 5.3.3.2 Method 0060 Train Accuracy and Precision Assessment

The quality control procedures that were implemented during the analyses of these samples for the purposes of assessing the general accuracy and precision of the analytical processes included the analysis of LCSs, LCSDs, PDS, and MS/MSD samples. Laboratory control samples test the accuracy and precision of the laboratory processes independent of stack gas matrix effects. The quantitative recovery of PDSs provides a second indicator of accuracy for the metals analysis for matrices (e.g., the front-half and back-half samples) from which matrix spikes cannot be prepared without affecting detection limits.

These spikes act as "internal standards" and signal when problems are encountered with the digestate matrix. When acceptable quantitative recoveries are observed for a PDS, the sample introduction onto the ICP is considered to be taking place correctly and the instrument has quantified the analytes present in the digestate correctly. Low recoveries typically indicate that viscosity or matrix interference effects may have been encountered. Matrix spikes (post sampling) of mercury were applied to the four back-half fraction samples. Accurate and precise recovery of the spiked mercury indicates that the entire analytical process, including preparation, is in control.

Matrix spikes of the ICP metals were not performed for the back-half composite sample matrix because this process would raise detection levels while adequate quality control information can be otherwise obtained. In terms of compliance, SW-846 Method 0060 does not require matrix spike recovery information for the evaluation of metals train samples, while SW-846 Method 6010B requires them to be performed for each sample batch or sample delivery group (SDG). The reasoning behind the lack of matrix spike requirements in the Method 0060 relates to the difficulty of dividing a front-half train portion, which contains the particulate filter, into three equal portions of the PM sample. Dividing the filter is a precarious operation. Particulate matter may not be evenly distributed on the filter, and cutting the filter to obtain equal portions of particulate material on each portion of the filter is difficult to execute. Analysis of MS/MSD samples for the front-half composite sample in not technically advisable. Particulate matter is the fraction of the captured stack gas sample that contributes typically the greatest level of metals to the Method 0060 samples.

Post digestion spikes were allowed in the QAPjP as an alternative QC measurement to replace MS/MSDs for both the front half and the back half train fractions. Post digestion spikes give an adequate demonstration of recovery for these samples, and are allowed by quality procedures sections of SW-846 methods for flame atomic absorption and graphite furnace atomic absorption. When coupled with laboratory control samples, and laboratory control sample duplicates, the quality of Method 0060 train sample analysis can be completely evaluated.

The LCS and PSDs corresponding to the front-half composite samples exhibited acceptable recoveries for all target metals. The recoveries of the metals, were within the target recovery range of 75% to 125%, with the exception of manganese and mercury. Mercury recoveries were high for Run 1, Run 2, Run 4 and the Final Acetone and Nitric Acid Probe Rinse. Manganese exhibited a low recovery only for Run 4.

The back-half sample fractions were also supported by laboratory control sample analysis and post digestion spike analysis. Post digestion spikes were analyzed for the back-half samples from Runs 1 and 2. The laboratory control sample results were accurate and reproducible, and indicate that the analytical processes were in control. All of the post digestion spike results were acceptable, but the mercury results have little meaning because the native levels of mercury in the samples were very high relative to the spike levels. Matrix spikes of the Run 1 back-half fractions for mercury of the impinger number 4 contents, 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> impinger contents, and the 8N HCl rinse fractions were successful. Matrix spike results for the back-half composite sample for Run 1 could not be calculated because the native level of mercury was too high, relative to the spike level.

In general, the metals laboratory control sample, post digestion spike, and matrix spike recoveries indicate acceptable performance and provide a strong indication that the analytical processes used were in control. A review of the calculated RPDs (presented in the final STL data package and archived for this project), obtained from the analysis of LCS, indicates that these analytical processes are highly reproducible.

The data were reviewed for evidence of interelement interference because ICP (AES) analysis is subject to interelement interference from "high" levels of a few analytes. Aluminum (Al) is the only element present in the NWCF ETS samples that is in a sufficiently high concentration to be considered an interfering influence on the results for target analytes. The aluminum concentration for the Run 3 front-half composite sample is a potential interferent that could increase the results for vanadium. There is no effect, because the result for vanadium for this sample was non-detectable.

## 5.4 Particulate Matter and Acid Gases

The standard EPA Method 0050 HCl/Cl<sub>2</sub>/Particulate Train (SW-846 Method 0050) configuration was used to collect samples from the NWCF ETS offgas for the assessment of PM and acid gas vapors. An offgas sample having a nominal volume of 3.0 dscm was isokinetically collected over 3-4 hours during each sampling run. A blank train and applicable reagent blanks were collected to support the QA/QC requirements specified in the QAPjP.

A final probe rinse sample was collected after the test runs for particulate analysis. The glass-lined sampling probe could not routinely be removed from the offgas sampling manifold for rinsing after each run for safty reasons as previously discussed. Instead, the front-half acetone rinse samples for particulate were limited to accessible tubing, filter housing elements and other connecting glassware that were installed outside the manifold and probe assembly. The glass-lined probe was only removed from the manifold after all offgas sampling trains were collected. An acetone probe rinse sample of the glass-lined probe was collected after the test runs to assess the maximum amount of particulate materials that adhered to the inside of the probe during testing. The probes were rinsed three times with acetone, and the samples were submitted for particulate analysis.

## 5.4.1 Acid Gas Target analytes

The target analytes for this method are hydrogen chloride (HCl), chlorine (Cl<sub>2</sub>), hydrogen fluoride (HF), nitrate (NO<sub>3</sub>-), nitrite (NO<sub>2</sub>-), and PM.

### 5.4.2 Analysis of PM and Acid Gases

The particulate samples collected included two fractions: (1) an acetone rinse of the probe and filter housing assembly, and (2) a particulate filter. Composite samples containing the contents of the first, second, and third impingers of the train were collected for HCl, HF, HNO<sub>3</sub>, and HNO<sub>2</sub> analysis. The first impinger was empty at the beginning of the sampling run and served as a moisture knockout impinger. The second and third impingers were each initially charged with 100 mL of 0.1N H<sub>2</sub>SO<sub>4</sub>. These acidic impingers allowed for the dissociation and collection of HCl, HF, HNO<sub>3</sub>, and HNO<sub>2</sub> from the offgas. A composite sample of the fourth and fifth impinger contents, which were each initially charged with 100 mL of 0.1N NaOH, were analyzed for Cl<sup>-</sup>, F<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, and NO<sub>2</sub><sup>-</sup>. Chlorine gas (Cl<sub>2</sub>) present in the offgas does not freely dissociate in the acidic H<sub>2</sub>SO<sub>4</sub> medium; therefore, it passes through the first three impingers as an unreacted gas and reacts when it comes in contact with the NaOH solution as follows:

$$Cl_2 + 2OH^- \xrightarrow{NaOH} 2OCl^- + Cl^- + H_2O$$

At STL, the particulate filter samples were dried at 105°C and the acetone probe rinse samples were evaporated to dryness at room temperature. Both fractions were stored in a dessicator for 24 hours, then analyzed gravimetrically. Replicate weights were obtained until constant weights were achieved.

The difference between pre-sampling and post-drying gravimetric measurements were then calculated for each sample.

The 0.1N H<sub>2</sub>SO<sub>4</sub> and 0.1N NaOH impinger samples were analyzed by ion chromatography using SW-846 Methods 9056 and 9057, as implemented by STL Analytical Laboratories Method, KNOX-WC-0005, *Anion Analysis by Ion Chromatography*, KNOX-WC-0005 (April 20,1999). The calibration range extended from 0.5 mg/L to 20 mg/L for all of the target anions. In order to quantify all of the target anions, several analyses were conducted at different dilution factors that ranged from 1 to 10. Dilution was performed both to bring the sample concentration within the instrument calibration range and to overcome matrix effects. The optimum value was chosen for reporting, with the lowest achieved detection limits reported in each case. The final data for each anion were based on analyses that were within the calibration range of the instrument.

The tabulated data summaries provided in the STL Final Report are listed in Appendix A. Each anion result is reduced to a "Risk Result" by selecting the default value for use in risk assessment calculations in accordance with project guidelines.

### 5.4.3 PM and Acid Gas Data Quality Assessment

Samples were shipped using Federal Express overnight delivery service to the laboratory. All Method 0050 train samples that were collected during the offgas sampling program were received at the laboratory in good condition. No losses of samples due to breakage or loss of shipment occurred. All Method 0050 train fractions were processed on schedule, as required by the QAPjP, and analytical results were obtained for all of the expected analyses.

Sufficient data have been collected and reviewed to address the relative precision and accuracy of the particulate and anion target analyte measurements. The data quality indicators present sufficient evidence that the data are of acceptable quality and are usable for the NWCF ETS emissions inventory.

## 5.4.3.1 Method 0050 Train Reagent Blank Assessment

A particulate filter, the  $0.1N\ H_2SO_4$  and  $1N\ NaOH$  reagents, and a sample of the INTEC supplied deionized (D.I.) water were collected during sample collection and were processed as reagent blanks in order to assess the presence of background analytes. These reagent blank samples were analyzed for the same parameters as the actual train samples, and showed minimal background levels of the target analytes. Chloride was detected in the  $0.1N\ H_2SO_4$  reagent blank at a level that was below the RL for the laboratory. Nitrate was detected in the  $0.1N\ NaOH$  reagent blank and the D.I. water reagent blank at low levels that had no impact on the final results. The sample results have been presented without blank corrections with the exception of tare subtraction required for the particulate analysis. All sample calculations of offgas concentrations were performed without blank or background correction.

## 5.4.3.2 Method 0050 Blank Train Assessment

A set of blank train samples was collected in conjunction with the four Method 0050 runs. The blank train samples exhibited similar results as the reagent blanks. There was no evidence of significant contributions to anion contributions in the sample results as a consequence of the sample train component preparation or handling.

## 5.4.3.3 Laboratory Control Sample and Matrix Spike Sample Assessment

Laboratory control sample percent recoveries indicate that the analytical process was in control. However, there were reduced recoveries of chloride in the matrix spike samples, due to apparent matrix effects. The other anions exhibit percent recoveries that are within target acceptance limits. Overall recovery of all of the target anions was sufficient to indicate that the data are useful for their intended purposes. The RPD results for both the LCS and the matrix spike samples indicate that a high level of precision is represented by this data.

## 6. PROCESS STREAM CHARACTERIZATION RESULTS

During the NWCF ETS off-gas emissions sampling activities, the NWCF ETS was being used to process a blend of two parts by volume of solution from INTEC Tank Farm Facility (TFF) vessel WM-184 and one part by volume of solution from INTEC TFF vessel WM-181. The campaign to process this blend was initiated on May 4, 2001 and continued through December 2001. At the same time the off-gas sampling was being performed, RCRA-quality liquid samples of NWCF ETS process streams were collected for analysis. These samples were collected under the protocols and QA/QC specified in sampling and analysis PLN-613 and PLN-407, the ALD QAPjP for environmental samples. These samples consisted of one each of the blended feed, condensed overheads, and concentrated bottoms. In addition, process samples of the feed were analyzed prior to initiation of processing the blend in the NWCF ETS to ensure that the chemistry of the feed solution was compatible with process equipment. Finally, samples of the condensed overheads and the concentrated bottoms from the first several batches processed were analyzed to ensure that the chemistry of those streams was compatible with down-stream process equipment. It should be noted that the non-RCRA samples were not analyzed per RCRA protocol; therefore limited QA/QC controls were evaluated.

These data obtained for the process streams may be used for component material balances around the NWCF ETS and are included in this report to provide a complete picture of the NWCF ETS and to provide a convenient location to obtain the data for subsequent system analyses. It should be noted that some limitations exist with this data. First, the INTEC Analytical Laboratory Department (ALD) that analyzed these samples has a more limited standard target analyte list for volatile organic compounds and semi-volatile organic compounds than the contract laboratory that analyzed the off-gas samples. The INTEC ALD TAL of volatile and semi-volatile organic compounds are contained in Appendix C. Second, some of the process samples (especially those taken to ensure compatibility with the NWCF ETS and down-stream equipment) were not taken at the exact same time as the off-gas samples. However, it is reasonable to assume that the process control system provides consistent batching of the feed streams and consistent control of the process variables. Third, the samples taken prior to initiation of the blend campaign, as well as those taken during processing the first several batches were only analyzed for a limited number of analytes. Since the purpose of these samples was to ensure compatibility between the solution chemistry and the process equipment, only those analytes that might challenge the envelopes of associated safety bases were targeted.

The results of the inorganic analyses of the feed samples are contained in Table 13, those results for the concentrated bottoms are contained in Table 14, and those for the condensed overheads are contained in Table 15. Organic compound analyses were only performed on one sample from each of the three streams; therefore, the organic analysis results for all three streams are contained in Tables 16 and 17. The INTEC ALD analytical reports for these samples are contained in Appendix C.

Table 13. Inorganic analyses of feed streams processed during NWCF ETS off-gas emissions sampling.

Analysis Log #:	0009272	0009274	0010022	00100415	0010164	0010167	0104103	0104125	0104142	0105062	0105063	0105112	0106071
Analysis, units:													
Sp. G.		1.1602	1.1549			1.1579	1.2220	1.2201	1.2409	1.1976			
Acid, N		1.564	1.596			1.633	1.842	1.823	1.854	1.781			1.711
Al, <i>M</i>		0.2209	0.2189			0.2127	0.526	0.537	0.521	0.451			0.42448
Sb, <i>M</i>													<0.000002
As, <i>M</i>													0.00000447
Ba, <i>M</i>													0.0000290
Be, <i>M</i>													0.0000105
B, <i>M</i>		0.0130	0.0134			0.0137	0.0076	0.0076	0.0082	0.00971			
Cd, <i>M</i>		0.00411	0.00426			0.00420	< 0.000948	< 0.000948	< 0.000948	0.00156			0.00156
Ca, <i>M</i>		0.04376	0.04526			0.04684	0.01618	0.0163	0.0169	0.02528			
Cl, M		0.0179	0.0117			0.0117	0.0276	0.0278	0.02835	0.0226			
Cr, <i>M</i>		0.00275	0.00252			0.00284	0.00169	0.0017	0.0018	0.00194			0.00196
Co, <i>M</i>													0.00000283
Cu, <i>M</i>													0.0004244
F, <i>M</i>		0.08485	0.1365			0.8122	0.0291	0.0258	0.0260	0.0500			0.04351
Fe, M		0.0113	0.0121			0.0129	0.01422	0.0138	0.0149	0.0133			
Pb, <i>M</i>													0.0005369
Mn, <i>M</i>													0.007841
Hg, <i>M</i>		0.00198	0.00102			0.000603	0.000748	0.000633	0.000678	0.000613			0.000743
Ni, <i>M</i>													0.00113
NO <sub>3</sub> , <i>M</i>		2.839	3.111			3.222	3.91	3.89	3.96	3.279			
PO <sub>4</sub> , <i>M</i>													
K, <i>M</i>		0.821	0.124			0.127	0.0895	0.0880	0.0880	0.0987			
Se, M													< 0.000003
Ag, M													0.0000011
Na, M		5.70	0.848			0.853	1.32	1.31	1.36	1.17			
SO₄, <i>M</i>		0.0316	0.0343			0.0385	0.0162	0.0087	0.0106	0.0288			
Tl, <i>M</i>													<0.000001
U, <i>M</i>			0.00032			0.00031	0.000185	0.000184	0.000168	0.000206			0.000209

Table 13. Inorganic analyses of feed streams processed during NWCF ETS off-gas emissions sampling.

Analysis Log #:	0009272	0009274	0010022	00100415	0010164	0010167	0104103	0104125	0104142	0105062	0105063	0105112	0106071
Analysis, units:													
V, <i>M</i>													0.000012
Zn, M													0.0006313
Zr, <i>M</i>		0.0053	0.0054			0.0053	< 0.00278	< 0.00278	< 0.00278	< 0.00279			
H-3, mCi/L	0.0158			0.0154	0.0126						0.0223	0.0219	
Co-57. mCi/L							0.01171						
Co-60, mCi/L		0.0705	0.0649			0.0627	0.01357	0.01710	0.01697				
Sr (total), mCi/L	20.4			24.28	20.69						6.76	16.7	
Cs-134, mCi/L		0.0705	0.0635			0.0646		0.00546	0.00555	0.0212			
Cs-137, mCi/L		25.8	26.1			26.2	12.8	14.24	14.38	19.4			
Eu-154, mCi/L		0.214	0.224			0.197	0.0299	0.0319	0.0324				
Eu-155, mCi/L		0.0484	0.0451			0.0451							
Nb-94, mCi/L		0.00316	0.00308			0.00373				0.00209			
Zr-95, mCi/L		0.0125	0.0115			0.0121				0.00477			
Sb-125, mCi/L						0.0368			0.130				
Am-241, mCi/L													
TIC,¹ μg/mL													<119
UDS, g/L													0.619
Total inorganic c	arbon.			-									

Table 14. Inorganic analy	yses of bottoms	streams during	NWCF ETS off-	gas emissions
Analysis Log #:	0105106	0106146	0106214	0106233
Analysis, units:				
Sp. G.	1.0747	1.3564		1.3036
Acid, N	2.559	2.868	1.741	2.830
A1, <i>M</i>	0.518	0.859	0.31256	0.711
Sb, <i>M</i>			0.0000128	
As, M			< 0.000008	
Ba, M			0.0000216	
Be, M			0.0000067	
B, M				
Cd, M			0.00131	
Ca, M				
Cl, <i>M</i>	0.0226	0.02866		0.03732
Cr, M			0.001674	
Co, M			0.0000241	
Cu, M			0.0003003	
F, <i>M</i>	0.0466	0.0900	0.052775	0.0684
Fe, <i>M</i>	0.0100	0.0700	0.00-2	
Pb, <i>M</i>			0.0004164	
Mn, <i>M</i>			0.006174	
Hg, <i>M</i>			0.000568	
Ni, M			0.0009426	
	4.03	5.27	0.00007420	7.22
$NO_3, M$	0.003871	0.01399		0.01382
$PO_4, M$	0.003871	0.129		0.154
K, <i>M</i>	0.0139	0.129	< 0.000012	0.154
Se, M			<0.000012	
Ag, M	0.505	1.56	<u> </u>	1.91
Na, M	0.505	1.30		1.91
SO <sub>4</sub> , M			<0.000004	
Tl, <i>M</i>	0.000047	0.000211	<0.000004	0.0004257
U, <i>M</i>	0.000247	0.000311	0.000261	0.0004357
V, <i>M</i>			0.000011	
Zn, M			0.000494	
Zr, M	0.0100	0.0151		
H-3, mCi/L	0.0108	0.0151		
Co-57, mCi/L				
Co-60, mCi/L	0.0352			
Sr (total), mCi/L				
Cs-134, mCi/L	0.0236			
Cs-137, mCi/L	21.2			
Eu-154, mCi/L	0.0904			
Eu-155, mCi/L				
Nb-94, mCi/L				
Zr-95, mCi/L	0.00528			
Sb-125, mCi/L				
Am-241, mCi/L				
TIC,¹ μg/mL			<119	
UDS, g/L	5.3	1.187	1.288	0.725
Total inorganic carbon.				

Table 15. Inor	ganic analys	ses of conde	nsate strean	ns during N	WCF ETS o	off-gas emis	sions
Analysis Log #:	0105061	0105073	0105087	0105092	0106145	0106221	0106241
Analysis, units:							
Sp. G.	1.0127	1.0121	1.0131	1.0131	1.0138		1.0130
Acid, N	0.467	0.450	0.476	0.480	0.498	0.514	0.471
Al, <i>M</i>	0.00108	< 0.0008	< 0.00042	< 0.00075	< 0.00042	0.00006312	< 0.00042
Sb, <i>M</i>						< 0.0000004	
As, <i>M</i>						< 0.0000004	
Ba, <i>M</i>						0.000000066	
Be, <i>M</i>						< 0.0000001	
B, <i>M</i>							
Cd, <i>M</i>						< 0.00000004	
Ca, <i>M</i>							
Cl, <i>M</i>	0.004928	0.00485	0.005156	0.00525	0.005602		0.00539
Cr, <i>M</i>	0.001,540	3,33,102				0.00000052	
Co, <i>M</i>						< 0.0000002	
Cu, <i>M</i>						< 0.0000003	
F, <i>M</i>	< 0.00041	< 0.00037	< 0.00037	< 0.00037	< 0.00018	0.0001791	< 0.00018
Fe, <i>M</i>	VO.00041	10.00057	-0.00057	10.00057	0.000.0	0.000.75	3133333
Pb, <i>M</i>						< 0.0000004	
Mn, <i>M</i>						0.00000024	
	0.0000229	0.0000189	0.0000186	0.0000115	0.0000274	0.00001969	0.0000163
Hg, M	0.0000229	0.0000169	0.0000100	0.0000115	0.0000274	0.00000055	0.0000105
Ni, <i>M</i> NO <sub>3</sub> , <i>M</i>	0.4011	0.4198	0.02049	0.4233	0.4393	0.00000055	0.4007
• .	0.4011	0.4196	0.02049	0.4255	0.4373		0.4007
$PO_4, M$							
K, <i>M</i>						< 0.0000007	
Se, M						<0.0000007	
Ag, M						<0.0000002	
Na, M	0.000153	0.000120	<0.00003	0.000113	0.000083		< 0.000059
$SO_4, M$	0.000153	0.000120	< 0.00003	0.000112	0.000083	< 0.0000002	<0.000039
Tl, <i>M</i>	-0.0000014	-0.0000014	-0.0000014	<0.0000014	<0.0000014		< 0.0000014
U, <i>M</i>	< 0.0000014	< 0.0000014	< 0.0000014	< 0.0000014	< 0.0000014	<0.0000014	<0.0000014
V, <i>M</i>				F		<0.0000002	
Zn, M						0.00000081	
Zr, M							
H-3, mCi/L				•			
Co-57, mCi/L							
Co-60, mCi/L							
Sr (total), mCi/L							
Cs-134, mCi/L							
Cs-137, mCi/L							
Eu-154, mCi/L							
Eu-155, mCi/L							
Nb-94, mCi/L							
Zr-95, mCi/L							
Sb-125, mCi/L							
Am-241, mCi/L							
TIC,¹ μg/mL						<23.8	
UDS, g/L	none visible	none visible	none visible	none visible	none visible	0.0	none visible
1. Total inorganic c	arbon.		· · · · · · · · · · · · · · · · · · ·				

Stream	Blended I		Botton		Condensate	
	Result	LQ	Result	LQ	Result	LQ
Analyte, units:						
TOC, μg/mL	608.462		754.99		147.853	
Chloromethane, μg/L	<10	U M	<10	U	<10	U
Vinyl Chloride, μg/L	<10	U	<10	U	<10	U
Bromomethane, μg/L	4	J	37	ВМ	160	EBM
Chloroethane, µg/L	<10	U	<10	U	<10	U
Trichlorofluoromethane, µg/L	<10	U	<10	U	<10	U
1,1-Dichloroethene, μg/L	<10	U	<10	U	<10	U
1,1,2-Trichloro-1,2,2-	<10	U	<10	U	<10	U
Carbon disulfide, µg/L	<10	U	<10	U	<10	U
Acetone, μg/L	9	J	<20	UΖ	32	Y
Methylene chloride	<10	U	<20	UΖ	<20	UΖ
Trans-1,2-dichloroethene, μg/L	< 10	U	<10	U	<10	U
1,1-Dichloroethane, µg/L	<10	U	<10	U	<10	U
Cis-1,2-dichloroethene, µg/L	<10	U	<10	U	<10	U
2-Butanone, μg/L	<10	U	<10	U	<10	U
Chloroform, µg/L	<10	U	<10	U	<10	U
1,1,1-Trichloroethane, µg/L	<10	U	<10	U	<10	U
Carbon tetrachloride, µg/L	<10	U	<10	U	<10	U
Benzene, µg/L	<10	U	<10	U	<10	U
1,2-Dichloroethane, µg/L	<10	U M	<10	U	<10	U
Trichloroethene, μg/L	<10	U	<10	U	<10	U
1,2-Dichloropropane, µg/L	<10	U	<10	U	<10	U
Bromodichloromethane, µg/L	<10	U	<10	U	<10	U
Cis-1,3-dichloropropene, µg/L	<10	U	<10	U	<10	U
4-Methyl-2-pentanone, μg/L	<10	U	<10	U	<10	U
Toluene, μg/L	<10	U	<10	U	<10	U
Trans-1,3-dichloropropene, μg/L	<10	U	<10	U	<10	U
1,1,2-Trichloroethane, µg/L	<10	U	<10	U	<10	U
Tetrachloroethene, μg/L	<10	U	<10	U	<10	U
2-Hexanone, μg/L	<10	U	<10	U	<10	U
Dibromochloromethane, µg/L	<10	U	<10	U	<10	U
Chlorobenzene, µg/L	<10	U	<10	U	<10	U
Ethylbenzene, µg/L	<10	U	<10	U	<10	U
M-xylene and p-xylene, μg/L	<20	U	<20	U	<20	U
O-xylene, μg/L	<10	U	<10	U	<10	U
Styrene, µg/L	<10	U	<10	U	<10	U
Bromoform, µg/L	<10	U	<10	U	<10	U
1,1,2,2-Tetrachloroethane, μg/L	<10	U	<10	U	<10	U
Tentatively Identified Compounds:						
Unknowns, number	1/10	J				
LQ = lab qualifiers (see Appendix C for de	efinitions)	<del></del>				

Stream	Blende	d Feed	Botto	ms	Condensate	
Analysis Log #:	0106	071	01062	221	01062	214
	Result	LQ	Result	LQ	Result	LQ
Analyte, units:						
TOC, μg/mL	608.462		754.99		147.853	
N-Nitrosodimethylamine	<20	U M	36		42	
Pyridine	<20	U	<20	U	<20	U
Phenol	<20	U	<20	U	<20	U
bis(2-Chloroethyl)ether	<20	U	<20	U	<20	U
2-Chlorophenol	<20	U	<20	U	<20	U
1,3-Dichlorobenzene	<20	U	<20	U	<20	U
1,4-Dichlorobenzene	<20	U	<20	U	<20	U
1,2-Dichlorobenzene	<20	U	<20	U	<20	U
2-Methylphenol	<20	U	<20	U	<20	U
bis(2-Chloroisopropyl)ether	<20	U	<20	U	<20	U
3 & 4-Methylphenol	<20	U	<20	U	<20	U
N-Nitroso-di-n-propylamine	<20	U	<20	U	<20	U
Hexachloroethane	<20	U	<20	U	<20	U
Nitrobenzene	<20	U M	<20	U	<20	U
Isophorone	<20	U	<20	U	<20	U
2-Nitrophenol	<20	U	<20	U	<20	U
2,4-Dimethylphenol	<20	U	<20	U	<20	U
bis(2-Chloroethoxy)methane)	<20	U	<20	U	<20	U
2,4-Dichlorophenol	<20	U	<20	U	<20	U
1,2,4-Trichlorobenzene	<20	U	<20	U	<20	U
Naphthalene	<20	U	<20	U	<20	U
4-Chloroaniline	<20	U	<20	U	<20	U
Hexachlorobutadiene	<20	U	<20	U	<20	U
4-Chloro-3-methylphenol	<20	U	<20	U	<20	U
2-Methylnaphthalene	<20	U	<20	U	<20	U
Hexachlorocyclopentadiene	<20	U	<20	U	<20	U
2,4,6-Trichlorophenol	<20	U	<20	U	<20	U
2,4,5-Trichlorophenol	<20	U	<20	U	<20	U
2-Chloronaphthalene	<20	U	<20	U	<20	U
2-Nitroaniline	<20	U	<20	U	<20	U
Dimethylphthalate	<20	U	<20	U	<20	U
2,6-Dinitrotoluene	<20	U	<20	U	<20	U
Acenaphthylene	<20	U	<20	U	<20	U
3-Nitroaniline	<20	U	<20	U	<20	U
Acenaphthene	<20	U	<20	U	<20	U

Table 17. SVOC analyses of NWCF ETS streams during NWCF ETS off-gas emissions Blended Feed Bottoms Condensate Stream 0106221 0106214 0106071 Analysis Log #: LQ LQ Result LQ Result Result Analyte, units: 44 Μ 110 M 2,4-Dinitrophenol 420 DMH 4-Nitrophenol U <20 U M <20 <20 U M Dibenzofuran <20 U <20 U <20 U 2,4-Dinitrotoluene U U <20 U <20 <20 Diethylphthalate <20 UM< 20 U M U M < 20 4-Chlorophenyl-phenylether U <20 U < 20 U <20 Fluorene U U <20 <20 U < 20 4-Nitroaniline U <20 U <20 U <20 4,6-Dinitro-2-methylphenol U M <20 U M <20 U M <20 N-Nitrosodiphenylamine U <20 U <20 U <20 U U Tri-n-butyl phosphate U U Azobenzene 4-Bromophenyl-phenylether <20 U U <20 U <20 Hexachlorobenzene <20 U <20 U <20 U Pentachlorophenol U <40 U <40 U <40 Phenanthrene U <20 U <20 U <20 Anthracene U <20 U <20 U <20 Carbazole U <20 U U <20 < 20 Di-n-butylphthalate <20 U M <20 U M <20 U Fluoranthene <20 U <20 U U < 20 Pyrene U <20 U <20 U <20 Butylbenzylphthalate < 20 U <20 U <20 U 3,3'-Dichlorobenzidine <20 U M < 20 U <20 U M Chrysene U <20 U <20 U <20 Benzo(a)anthracene U U <20 U <20 <20 bis(2-Ethylhexyl)phthalate U <140 U <20 U <20 Di-n-octylphthalate U U <20 U <20 <20 Benzo(b)fluoranthene <20 U <20 U <20 U Benzo(k)fluoranthene U < 20 U <20 U <20 Benzo(a)pyrene U <20 U <20 U <20 Indeno(1,2,3-cd)pyrene U <20 U <20 <20 U Dibenzo(a,h)anthracene U <20 U <20 U <20 Benzo(g,h,I)perylene U <20 U <20 U <20 Tentatively Identified Compounds: 19/509 J 5/130 J Unknowns, number identified/conc. 7/501 J 17 J Tri-n-butyl phosphate

LQ = lab qualifiers (see Appendix C for definitions)

## 7. PROCESS OPERATING CONDITIONS

The NWCF ETS is operated as described in Section 2. The evaporator is initially filled with fresh waste solution. The temperature in the evaporator flash column, prior to initiation of stream is usually between 50 and 60°C. Steam flow is ramped from 0 to 1730 lb/hr in 15 to 45 minutes and then maintained for eight to ten hours until the desired solution density is obtained. Approximately 2 to 3 hours is required for the column to reach a full boil at around 100°C. Fresh feed solution is added to the evaporator until near the end of the batch. The temperature increases 3-4° throughout the batch as the concentration increases to the target density.

Offgas sampling at the beginning of the evaporator run was synchronized with the initiation of steam to the evaporator. Approximately 3 hours was needed to collect the offgas samples, including the SMVOC runs. This provided a representative average of the emissions during the startup period. SMVOC samples were actually started 15-20 minutes prior to steam initiation in order to capture any burst of volatile organic emissions on the onset of solution heating. Sample collection at the end of the evaporator batches was coordinated with the ETS operators in order to sample during the final 3-4 hours of the evaporator batch.

Process parameter data were collected during the NWCF ETS emissions sampling by the NWCF Distributive Control System (DCS). A history of key process variables was collected using fifteen-minute average data. The data were then tabulated for the times when the sampling was taking place. Appendix D compiles the process parameters for the sixteen sample collection runs.

The evaporator operated within normal operating parameters throughout the period of sample collection. The feed batches had very similar densities and temperatures, indicating very similar compositions. The total flow from the NWCF (containing the NWCF ETS emissions) was essentially constant limiting variation due to deposition or re-entrainment.

During the collection of sample train 0060-STRT-02, sampling was interrupted for an emergency drill while the evaporator batch continued. Sampling was interrupted from 0850 to 0922 on June 6, 2001 until sampling personnel were notified that they could continue. During this time, the average temperature in the evaporator increased from 76.4 to 97.4 degrees.

## 8. DQO ASSESSMENT AND PROJECT SURVEILLANCE

Data quality objectives (DQOs) for the NWCF ETS offgas emissions project are defined in the quality assessment project plan (QAPjP, company document PLN-879). Sample collection in the field was coordinated by the Project Technical Leads (PTL) with independent surveillance performed by the Project Quality Assurance Officer (PQAO). The sample collection activities were monitored by the PTL and PQAO, thus ensuring that the sample collection activities were completed in accordance with the test plan and QAPjP and that the samples were maintained under proper custody and conditions at all times. All changes to the test plan required advance approval from the PTL and PQAO prior to being implemented during sample collection. A standard field change form was used to document the approvals for these changes.

The services of the INEEL Sample Management Office (SMO) were not enlisted to review the analytical data. This was previously completed on the NWCF Calciner offgas emissions inventory, but was not within the budget constraints of the current project. Therefore, a cursory review of the analytical data QA/QC requirements was completed by the PQAO and is provided in lieu of the Limitations and Validation (L&V) reports that are provided by the SMO on previous projects.

# 8.1 Documented Field Changes

The QAPjP allowed for in-field changes to requirements of the QAPjP and sampling protocols as long as such changes were approved per Section 13.1 of the QAPjP. Also allowed by the QAPjP were properly approved changes to the sampling checklists. Seven field change forms capturing 11 different requested variances from the QAPjP or standard protocol were approved.

### 8.1.1 VOC Sample Collection

Method 0031 for VOCs still requires the storage of Tenax® and Anasorb-747® tubes at less than 10°C after tube conditioning, during transport, and up to the time of tube usage in sampling. This is a typical protocol deviation among current laboratory service providers, since it is deemed unnecessary in conjunction with the common practice of sealing the resin tubes in air-tight containers and using a trip blank to identify potential fugitive contamination that may occur prior to use of the tubes in the field and during their return to the laboratory. Conditioned sample media availability, schedule slippage, and laboratory technical guidance were critical factors in the project field decision to waive this requirement via a field change once it was determined that the laboratory had not complied.

The method requirement that conditioned media tubes not be exposed to severe pressure variations during transport is satisfied amply by the multiple layers of containment used by the contract laboratory providing the sample tubes. Therefore, in this regard, no field change was applicable. It should also be noted that all of the Tenax® and Anasorb-747® tubes were cooled to 0-10°C immediately following sample collection.

The initial intent of the contract sampling team was to not preserve aqueous samples collected for VOCs (Method 0031) by acidification with 8N HCl to a measured pH of < 2. The BBWI Project Technical Lead and the PQAO pointed out that it would be preferable to perform this standard EPA preservation protocol on such aqueous samples and on associated reagent water blanks. A field change to the sampling checklist reflecting that guidance was approved and sampling team was so instructed.

Another field change was proposed and approved to require another pair of Tenax®/Anasorb-747® tubes be collected as a field blank for the second day of Method 0031 sampling. The project had already varied in the QAPjP from the standard method requirement for a field blank every two hours of sampling based on the expected sampling period and reviews of cost /benefit and technical applicability of this QC-related method requirement to sample train operations, as scheduled.

The last field change related to the 0031 sample trains and protocols was one to keep the flowrates at or below 0.5 Lpm rather than going with the standard EPA Method 0031 upper limit of 1 Lpm. This allowed four sets of tubes to be ran in the 0031 sampling trains over a typical sampling interval meeting or exceeding the method target sampling time of 2 hours. This is compliant with the intent of Section 1.8 of the standard EPA SMVOC method.

## 8.1.2 SVOC Sample Collection

One field change form was approved specific to the Method 010 trains for SVOCs. This change incorporated a final field dilution of the condensate fractions from these trains from just under a total volume of 500 mL to a total volume of 1 liter. This reflects a dilution normally performed on the samples after receipt at the laboratory. By performing this dilution in the field under close supervision of the project tech lead and sample team leads, the project was readily able to comply with the 70 Bq/gram ceiling limit for shipping the associated SVOC samples in a non-radiological classification per DOT regulations.

### 8.1.3 SCS Trains and Screening

A field change was approved to allow for cancellation of a scheduled third radiological contamination evaluation train (SCS-EVAP-3), based on consistent and low radioactivity results obtained for all previous sample contamination trains and routine gamma counting screening that was performed in accordance with MCP-1173, *Package and Ship NWCF Offgas Emissions Samples Offsite for Analysis*, Revision 2.

## 8.1.4 Metals and Anions Sample Collection

No field changes were required for the metals and anions sample collection runs. All runs proceeded in accordance with the sample collection checklists and test plans.

### 8.1.5 Miscellaneous Decisions

A final field change capturing five different field decisions was processed and approved. Per QAPjP language, not all of the decisions under this field change were required by project planning documents to be documented by a field change. However, the PQAO and the PTL decided it would be a best management practice to well document these decisions, and this process was the most accommodating and represented a configuration management control mechanism totally internal to the project. The following change elements were approved on this change form.

## Element1:

It was determined that the pH meters supplied by the sampling contractor and in the field for this project could not meet reasonable time to stabilization criteria. Based on this fact, a field decision was made by PQAO and PTL to allow the samplers to do all pH measurements with pH paper versus requiring a meter. This was facilitated by the fact that in-field neutralizations of liquid sample fractions was not necessary as in the emissions inventory for

the NWCF Calciner. Approval was given at beginning of sample collections for the use of indicator paper for pH measurements in NaOH impinger sample fractions due to the harsh effects of the high pH.

#### Element 2:

This documents and certifies the assumption made early in project that the oxygen monitor readings of the sampled gas stream basically reflected the concentration of oxygen in ambient air. This assumption was used to relax standard EPA method calibration protocols and was already allowed by the QAPjP.

#### Element 3:

In conjunction with Element 2 above, it was decided to use oxygen cylinders already in inventory and marked as 20.8% O<sub>2</sub> rather than secure standards as cited in 40 CFR 60. In an associated decision, it was determined that for this project, the percent drift determinations would be made over time lines comparable to the actual train run times versus the 24-hour comparison baselines found in the 40 CFR 60 regulations which were inferentially tied to continuous monitoring.

#### Element 4:

Similar to Element 3 above, two other decisions related to the operation of the "continuous emissions monitoring system (CEMS)" monitors were made before inception of sample collections. First, use of a non-heated Perma-Pure® dryer to condition sample stream was authorized. Second, a rotameter with a 0-0.8 Lpm range was selected for use with the  $O_2$  monitor. This resulted in a target flowrate of 0.75 Lpm rather than the 1.0 Lpm cited in the standard method.

### Element 5:

Typical condensate recoveries from Method 0050 and Method 0060 trains were much less than 1.0 ml. The condensate and knockout (KO) rinse was added to impinger K1- K2 contents and rinses. This was per approved checklists. Additionally, for VOCs, the condensate fraction was topped to fill a 20-ml standard VOA vial and acidified with 8N HCl to a pH of <2. Reagent water blanks for VOCs analyses were processed in same fashion as 0031 condensates. In retrospect this decision element is redundant to other field changes already processed.

# 8.2 Data Quality Indicators

As of this point, all post-laboratory data evaluations for achievement of qualitative objectives and for quantitative data quality indicator acceptance criteria have been performed by the PTL and the PQAO. With the exception of a final evaluation of inter-train precision based on calculations of RSDs associated with results of surrogates from all trains for VOCs and all trains for SVOCs, the data quality indicators presented in the QAPjP have all been evaluated.

Reviews of analytical reports indicated that method performance and associated QC, as depicted in the lab reports, met analytical method and project planning document requirements, with a few instances of failures to meet individual QC criteria. The results for the vast majority of

associated QC data meet QAPjP criteria and support current designated project uses of this data. As stated, an independent analytical data validation to the cited Statement of Work (SOW) requirements found in ER-SOW-156, INEL Sample Management Office (SMO) Statement of Work for Inorganic & Miscellaneous Classical Analyses, Revision 1, ER-SOW-169, Statement of Work for Organic Analyses Performed for INEL Sample Management Office, Revision 0, or in associated standard analytical methods has not been performed. A full data validation process could potentially lead to additional data qualifications based on standard methods or cited SOWs.

## 8.3 Sampling Documentation Reviews

As in past emissions tests conducted for the NWCF Calciner, all field sheets related to sample collections were reviewed in the field as the sampling proceeded. Reviews were conducted by both the PTL and by the PQAO, and were completed to schedule per the QAPjP. Minor omissions or errors in field-level paperwork were therefore immediately caught and corrections were implemented while the sampling team was still in the field, and/or, if deemed necessary, so that re-sampling could occur with minimal delays or additional costs. Most observations that required corrective actions involved the accuracy or completeness of field data forms, and these instances were actually infrequent with respect to the sheer volume of field data entries required for the project.

All data were manually recorded in the field into the associated logbooks, sample data sheets and sampling checklists associated for each sample train type used during project sample collections. The data on sample data sheets were then uploaded into laptop computer templates of these same respective forms. There were some observed anomalies between some of the field definitions on the paper copies versus those on the electronic templates. These were noted in the individual internal PQAO reports to the PTL during each train type (0031, 0030, 0050, 0060) testing period. The sampling contractor (SAIC) made accommodations in each case by either giving more specific instructions to the sample collectors regarding the required entries on the paper forms, or by changing appropriate entry fields in the electronic templates. All changes were reviewed and approved in the field as required by the QAPjP.

As a result of reviews of the field-level documentation, it was discovered that the sample collectors had noted that sample #3411, Set#1 Anasorb-747® tube for Train 0031-END-1 had a crack at one end. Subsequent reviews of the laboratory entries into the related chain-of-custody forms did not indicate that this condition was noted by lab personnel on receipt of the samples. The laboratory analytical results appear consistent with other corresponding Anasorb-747® results and meet the necessary surrogate and internal standard recoveries.

Reviews of calibration documentation for associated sampling equipment identified no deficiencies with respect to requirements. Reviews of chain-of-custody and request for analysis forms identified no deficiencies in those documents.

# 8.4 Records Management

Records associated with this project have been retained and filed in an approved secure central file location (CFL) per company records management requirements. Project records have been categorized and dispositioned as environmental records and are currently assigned a permanent retention period. Per the project QAPjP and related company requirements and procedures for the designation and management of quality records, the following records have been further characterized as quality records:

- Quality Assurance Project Plan (QAPjP)
- Test Plans
- Logbooks
- Certificates of Analysis
- Calibrations
- Field Data Sheets
- Field Changes
- COC/RFA
- Analytical Data and Emissions Calculations Spreadsheets
- QA Reports
- Limitation and Validation Reports (not applicable yet for this project phase)
- Final Reports

Some of the above records are in paper form, some in electronic format, and some in dual media. All are stored appropriately in locked cabinets and controlled key access within the designated CFL. A file index has been prepared for these project records, and is available at the CFL. All records in CFL have been assigned appropriate INEEL uniform file codes in compliance with associated company procedures for management of files and records.

# 8.5 Review of Spreadsheet Calculations

As part of their contracted services, the sampling contractor provided spreadsheets which captured all appropriate field sampling data, analytical data, and project-required plant operations data, and which calculated from this data estimated total emissions rates from the NWCF ETS. As referred to elsewhere in this report, these estimates are conservative contributors to the emissions related to the operation of the ETS itself because other sources of miscellaneous plant tank operations continuously emit purge and/or vessel sparging gases to this same NWCF offgas line.

The first line of independent review and QC of these contractor spreadsheets occurred internal to the contractor. A second 100% review was conducted by the PTL, with any required corrections being implemented in conjunction the contractor for sake of configuration management and documenting general agreement with changes. Lastly, the PQAO reviewed in excess of 20% of the resultant pertinent entries after each iteration of changes. This comprehensive and iterated approach exceeded the basic requirement of the QAPjP that the PQAO verify only 20% of all entries and resultant calculation values. Once the sampling contractor lead, PTL, and PQAO were in agreement with the acceptability of the spreadsheets, the spreadsheets were noted as verified and released to be used on final report compilations.

# 8.6 Analytical Results

## 8.6.1 Data Reporting and Flagging

The QAPjP, associated task order statements of work, and technical lead guidance to the analytical laboratory defined project-specific requirements for data flagging, assignment of "<" symbols, and selection of most conservative and technically defensible result values for purposes of inserting conservative (high-biased) estimators of emissions rates for each identified compound into the emissions calculations. These requirements were in addition to the standard method-related data qualification flagging, and derive from the following EPA guidance documents:

- EPA 1998a, Guidance on Collection of Emissions Data to Support Site-Specific Risk Assessments at Hazardous Waste Combustion Facilities, EPA530-D-98-002, August.
- EPA 1998b, Human Health Risk Assessment Protocol for Hazardous Waste Combustion Facilities, EPA530-D-98-001A, July, Appendix A-1, Table A-1.
- Rule 1. When there was a non-detect below the MDL in a train fraction for a given compound, then the lesser of the RDL or the RL was used.
- Rule 2. When the laboratory RL is greater than the RDL, and a compound was detected above the MDL in a train fraction, but the result was less than the RDL, then the RDL was used.
- Rule 3. When the RL is less than the RDL, and a compound was detected in a given train fraction above both the MDL and RL values, but less than the RDL, then the RDL was used.
- Rule 4. When the RL is less than the RDL, and a compound was detected above the MDL, but result was less than the RL, then the RL was used (not the RDL).
- Rule 5. Any compound that was detected above the RDL was used for the risk calculation applications and no "<" ("less than") sign was assigned for that result in the respective train fraction.
- Rule 6. When a result for a SVOC or VOC target compound in a given train fraction was greater than MDL, but less than the RL, and result was assigned an estimated flag, the "<" flag was also assigned.
- Rule 7. Whenever a "<" flag was assigned to any given train fraction result, then the "<" flag propagated to the train total for that compound, unless it was dropped due to protocol of rounding to significant figures in the train total summation process. (ASTM E29-93a 1999).

Rules 2 and 5 default to the RL, not the RDL, when the RL is less than the RDL, and the "hit" was less than the RL. This was justified since the RL is a statistically established conservative RL, established by the analytical laboratory. Therefore, team assessed uses of RL value for risk calculations as a justified approach.

Most of the lab-assigned data qualifier flags are based on lab methods and procedures, and are standard to a large majority of environmental services laboratories. For metals analyses, the QAPjP required "B" flags to be assigned to metals results greater than MDL but less than the Reporting Limit.

Additionally, a system of assigning project flags "P", "N," and "A" to train total results was developed to evaluate the significance of each given target analyte result for the complete sampling train based on the relative occurrence of real hits for the various train fractions that comprised each sample train. The "P" indicated that related compound was detected in some train fractions, but not all. The "N" indicated that there was no positive detection in any train fraction for that compound. Lastly, the "A" flag on the train total result indicated that the compound was detected in all of the train sample fractions.

#### 8.6.2 Blank Corrections

Blank corrections for metals trains are allowed by the cited EPA guidance documents and standard air sampling methods. Reagent blank corrections were only made where actual hits occurred in the reagent blanks.

The performance of the SMVOC blank sample indicated that residual acetone and methylene chloride was present in the sample collection line connecting the probe to the train setup. The resulted from failure to adequately purge the sample collection line following collection of the SVOC samples which involves rinsing the line with a mixture of acetone and methylene chloride. The VOC results showed a rapid decay of acetone and methylene chloride in successive sample fractions. In spite of the fugitive contamination (which was included in the emissions inventory), volatile species emissions were very low; therefore, this problem is not considered a significant limitation of the data set. In order to correct this anomaly, it is recommended that a separate VOST probe always be used to collect the SMVOC trains. In the present inventory, the Project Technical Lead authorized the SVOC Method 5 probe for the SMVOC runs to reduce the risks associated with insertion and removal of a second probe into the NWCF offgas duct.

### 8.6.3 Data Reporting

Data reporting from laboratory was comprised of Certificates of Analysis, EDDs, and actual final reports which included complete data packages constructed to the lab's data package level equivalent to the INEEL Tier 1 reporting requirements. The INEEL internal documents that prescribe these data package expectations are ER SOWs -156 and -169. These reporting requirements were meant to allow for later validation of data should project management decide to subject this data to either a Level A or a Level B data validation per existing INEEL Sample Management Office procedures. Such a future validation decision might be made based on further identified uses of data or other criteria such as permit support, operational changes, planned facility modifications, etc.

## 8.6.4 Analytical QC

### 8.6.4.1 MS/MSD and LCS/LCSD Samples

Although analytical performance is treated in Section 5 of this report, there are some laboratory related QC aspects of results which need to be discussed in this section. The first QC topic area deriving from QAPjP relates to the selection and analysis of matrix spike samples and matrix spike duplicate samples, as well as the alternative approach of analyzing laboratory control sample spikes and duplicates.

The structural configuration of these EPA standard methods air sampling trains and non-homogeneity of matrices present in the various train sample fractions collected from the trains make traditional application of matrix spikes, matrix spike duplicates, and even sample splits very difficult. Splitting of sample fractions for analyses to calculate precision estimators obviously raises associated detection limits for those trains. Simultaneous or sequential operation and sample collections from two or three complete trains in order to derive estimates of precision and accuracy can quickly become very expensive and time consuming.

For this project, the PTL and contract laboratory project manager determined that either MS/MSDs or LCS/LCSDs with surrogate spikes would be utilized by analytical lab for various analyses. These associated surrogate spikes and acceptance criteria are listed in Table 3-1 of QAPjP. The RPD and percent recovery of these spikes were determined and provided by the contract laboratory. Evaluation of these data quality indicators was then accomplished later by the project team, using the criteria found in QAPjP Table 3-1.

The approach taken by laboratory for SVOCs was to analyze a front-half Composite LCS/LCSD and a back-half composite LCS/LCSD. For the aqueous matrices associated with 0010 trains, not only was a LCS/LCSD pair analyzed, but the lab also performed a 3-way split of the condensate and impinger contents for Run 2, using two of the split fractions to develop a MS/MSD pair. This allowed for generation of important matrix-specific information, but also tripled the detection limits for the non-QC sample aliquot. This is also discussed to a degree in Section 5.

Additionally, the XAD-2® tubes utilized in SVOC sampling were pre-spiked with 200  $\mu$ g of a  $^{13}$ C<sub>3</sub>-Naphthalene sampling surrogate. The determined recoveries of this surrogate were evaluated later in the project, and are discussed in Section 5 of this report.. Application of this labeled surrogate occurs prior to sample train operation and is a comprehensive estimator for the overall accuracy of surrogate application, collection method, laboratory sample prep, and analytical method. Additionally, recovery result is an indicator of potential losses of surrogate (or other SVOCs) or sample media cross-contamination occurring during shipments of same media to field or during shipment of collected samples back to the lab. All of the recovery results for this labeled sampling surrogate were in control.

With regard to VOC analyses of 0031 train samples, the same approach was taken, except that the QAPjP listed the recovery surrogates typical to Method 0031 and the following matrix spike compounds: 1,1-dichloroethane, trichloroethane, benzene, toulene, and chlorobenzene. These were selected with technical guidance from the INEEL SMO and reflect a subset of the standard surrogates used in the laboratory method which implements SW-846 Method 8260B. No spikes of the Tenax® media prior to sample collection were required. Again, acceptance criteria for these data quality indicators related to precision in accuracy were presented in QAPjP Table 3-1. The analytical report from the lab does not discuss results for such a LCS/LCSD. This may be a potential project-specific deficiency in the VOCs analyses unless additional data are located in the raw data packages.

For the Method 0060 metals trains, performance of post digestion spikes in accordance with EPA Method 6010B was an authorized approach. Additionally, for mercury, the QAPjP specified a MS/MSD pair. All associated acceptance criteria in terms of RPDs and percent recoveries were given in QAPjP Table 3-1. Again, for the metals train configuration there is no technically representative way to pre-spike train fractions before sample collection. As of time of this report, PQOA has not confirmed that lab reported associated LCS/LCSD results required by QAPjP.

Given the complexities of these matrices, the project team chose not to define sample fraction selection criteria or required frequencies of MS/MSDs to the lab in the QAPjP. Ongoing technical

consultations between the analytical lab and PTL determined the exact approaches which were taken in this regard for MS/MSD and LCS/LCSD analyses. They are included in the Requests-for-Analysis that were included in the Task Order Specific (TOS) Statement of Work (SOW) for sample analysis.

## 8.6.4.2 Performance for Internal Standards and Surrogates

Results for internal standards and surrogates for the VOCs analyses appeared to be acceptable based on QAPjP criteria. One outlier was Anasorb tube sample #3364 where results were non-usable. Some difficulties were encountered in recovery of the internal standard perylene-d<sub>12</sub>, one of the six internal standards for the 8270C analysis of the SVOC samples. In order to obtain acceptable recovery of this standard, it was necessary for the analytical laboratory to dilute the samples. This increased the detection limits for the reference target analyte species. A more complete discussion on this anomaly is provided in Section 5 and the STL Final Analytical Report (STL 2001).

In summary, all QA/QC criteria meet the data quality objectives with only two notable exceptions. First, there was some difficulty in recovering perylene-d<sub>12</sub>, one of the six internal standards used by the contract analytical laboratory for the Method 8270C SVOC analysis. The project used a trial XAD-2<sup>®</sup> to determine that the performance based QA/QC indicators would likely be achieved without modifications to the sample collection or analytical procedures. Thus, the poor performance of perylene-d<sub>12</sub> was not expected. It was necessary to dilute the final sample volume using methylene chloride (the same organic solvent that is used to extract the samples from the sample collection media) and then to "re-shoot" the sample with the GC/MS instrument. Dilutions of 10-100 times were needed to achieve acceptable recoveries of the perylene-d<sub>12</sub> standard. The implication of this result is that a few of target SVOC analyte data can only be considered an estimate, although the data are still usable for the emission inventory. This should not be considered a serious limitation of the data since all of the SVOC target analytes were typically less than the laboratory RL. In order to avoid this problem in future evaporator emissions testing, it is recommended that successive 1 mL samples be withdrawn from the sample solution during concentration (i.e., "blowdown") of the methylene chloride extraction solvent. This should be completed for the first run to determine the maximum concentration that is possible without failing to meet the specified internal standards recovery efficiencies.

Second, surrogate compound recoveries for one Anasorb<sup>®</sup> tube (A-3364 in Run 0031-STRT-1) failed to meet the acceptable recovery range. The results of the three corresponding Anasorb<sup>®</sup> tubes for this run were averaged and substituted for this tube. Surrogate performance and internal standard performance for all other Tenax<sup>®</sup> and Anasorb<sup>®</sup> tubes was generally excellent; therefore, the quality of the four SMVOC runs provide an accurate measurement of the target VOC analytes. Thus, this limitation did not significantly impact the run results. The run total was comparable to the results for the other 3 SMVOC runs.

# 8.7 Request for Analysis and Chain-of-Custody Forms

The analytical services laboratory used for this project utilized a system of pre-printed labels for samples based on a predetermined master sample list, in conjunction with "Request-for-Analysis" (RFA) and COC forms. This system greatly minimized chances for sample identification errors during the sample collection process. This is critical when there are multiple sample fractions for each train, many of which must be accurately combined either in the field or at the laboratory after sample receipt and log-in.

Reviews of closed out COC forms indicated only one instance of receipt of a shipping container without all container seals being intact. That occurrence was for the final probes rinses collected on June 25, 2001. There were otherwise no indications of any sample abnormalities observed for any sample receipts by the lab. All shipments of samples for analysis for organics arrived within acceptable temperature ranges per EPA criteria for sample preservation. Per the standard EPA methods, samples from 0050 and 0060 trains were not cooled during or prior to shipment.

## 8.8 Field Assessments by PQAO

In adherence with the QAPjP requirements for field assessments of sampling activities, surveillances/assessments were conducted for a single complete train run from each type of sample train used during the testing period for the NWCF ETS emissions. As a result some field changes processed, and corrective actions were taken in the field whenever necessary. There was no necessity per company procedures to initiate any forms for potential discrepancies or nonconformance reporting. Corrective actions were facilitated in the field during the respective sampling period.

Multiple types of checklists were utilized. Some criteria were based on the sampling protocol checklists themselves, while others dealt with good lab practice and work-site housekeeping. Housekeeping and lab practices ranged between acceptable to exemplary. These aspects were very important to this project given the spatial constraints of the sample collections area, rapidity of work schedule, and the multiple tasks occurring within the associated work hoods. Importance ranged from quality to safety, spill prevention, waste management, and effective radiological controls.

Field observations by the PQAO of the sampling team use of pH meters supplied by their company to perform pH measurements on project samples exhibited a failure of these meters to perform adequately with regard to stabilized readings in the buffered calibration standards or project samples. The PQAO, PTL, and SAIC sampling team lead agreed that substitution of pH indicator paper for all field pH measurements had adequate accuracy. This was facilitated by the fact that no sample neutralizations were required for this project phase. The section of this report dealing with field changes also discusses these points.

For this project, it was technically determined prior to inception of field activities that CEMS was not required to monitor the same miscellaneous gaseous emissions components that were a concern when the NWCF Calciner was running and being tested for emissions. In the past these emissions parameters included O<sub>2</sub>, CO<sub>2</sub>, CO, NO, NO<sub>2</sub>, HCl, CH<sub>4</sub>, SO<sub>2</sub> and total hydrocarbons (THC) and these derived from the aspects of feed to the Calciner and the physical presence of combustion products in the NWCF Calciner process. HCl emissions of ETS were evaluated using the results of the samples collected from the scheduled Method 0050 trains. For this project, dynamic monitoring was considered to be of technical value only for O<sub>2</sub> emissions. Additionally, it was predicted that the oxygen levels would be extremely close to those of ambient air. The monitored results from O<sub>2</sub> monitor throughout sampling campaign did actually reflect that oxygen concentration levels in the sampled offgas were essentially the same as ambient air concentrations of O<sub>2</sub>.

Latitude was built into the QAPjP regarding the operation and the calibration of the  $O_2$  monitor. Relief was given from the regulatory (40 CFR 60) requirements for the associated calibration gases, % Drift measurements, calibration frequency, etc. Additionally, the technical applications to associating the  $O_2$  levels with fairly short interval sample train runs, versus the usual regulatory application of 24-hr continuous monitoring, justified the relaxed protocol. It was determined that an appropriate calibration frequency for this monitor was that the calibrations must

occur prior to and after each test period, but not to exceed 24 hours between calibrations. Requirements were adequately implemented in the field. As part of the assessment of the operation and calibrations of  $O_2$  monitor it was documented that samplers were misinterpreting one entry associated with the documenting of % Drifts for this monitor. They were appropriately instructed in correct completion of the calibration sheets, and no further problems were observed.

## 9. OFFGAS EMISSIONS AND HEALTH RISK

Species emissions rates were calculated for all target analytes and tentatively identified compounds emitted from the NWCF Evaporator Tank System. Data for the train totals listed in Appendix A and the field sample collection data listed in Appendix B were compiled in an Microsoft Excel Program spreadsheet to compute both emissions rates [g/s basis] and offgas concentrations [ $\mu$ g/dscm basis], relative to conditions in the NWCF offgas duct where sampling was performed. Both the emissions rates and offgas concentrations summary sheets are included in Appendix B. Only a limited interpretation of emissions trends and potential risk to public health has been made at this time.

## 9.1 Emissions Rates and Trends

It was postulated that the release of organic compounds and volatile mercury present in the tank wastes would be higher at the start of the batch when the evaporator is filled with fresh feed. Conversely, it was hypothesized that metals emissions rates would increase with the density of the evaporator contents since the mechanism for non-volatile metals is primarily attributed to aerosol entrainment. Evolution of the organics, however, especially the semi-volatile organics, is a function of the evaporator temperature.

Figure 6 shows the average evaporator vessel temperature during a typical Method 5 sample collection period at the start and the end of an evaporator batch. The temperature was initially lower at the start of the run when steam to the heating coils was initiated. The temperature gradually increased, reaching the desired operating temperature of approximately 100°C at around 130 minutes-almost 3/4 through the first sampling period. Fresh feed to the evaporator was not increased significantly until the evaporator temperature reached the boiling temperature. The sample collected at the end of the evaporator run was performed when the evaporator vessel was at the peak temperature. At this time, feed was continually being supplied to maintain a constant volume in the evaporator. The volume of waste solution fed to the evaporator during the final three hours of the batch varied, but was approximately equal to ½ of the evaporator batch.

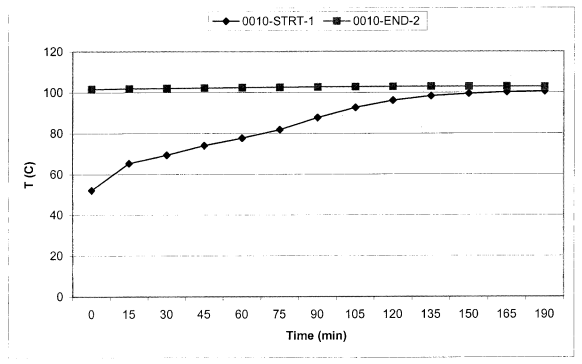


Figure 6. Average evaporator vessel temperature comparison for SVOC runs at the start and end on the evaporator batch.

## 9.1.1 Organic Compounds

Figure 7 plots the concentration of the 20 highest VOC compounds emitted from the evaporator. There are surprisingly small differences in the emissions rates at the beginning and end of the evaporator batch. The single highest volatile organic detected was dodecane, which was not a target analyte, but was reported as a tentatively identified compound. On a volumetric basis, the concentration of dodecane was only 54 ppbv. Acetone emissions were also relatively high, but still were in the low parts-per-billion range (*i.e.*, 30 ppb maximum). Some of the other volatile organic measurements were actually higher at the end of the batch, although the results were near the method detection limits and were susceptible to some process variations and sampling uncertainty at these lower levels.

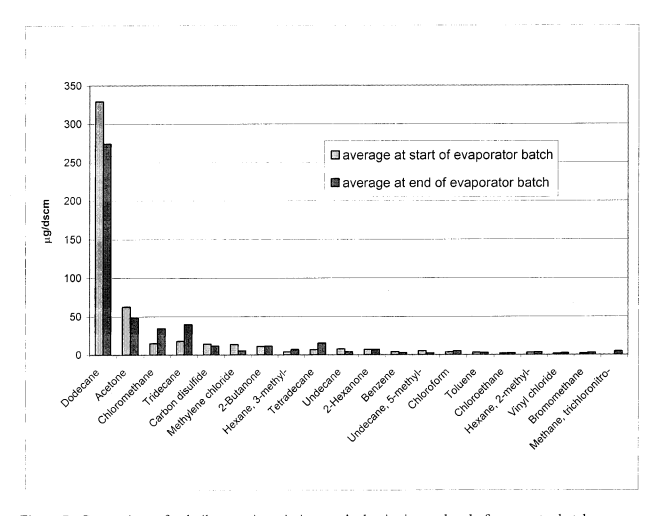


Figure 7. Comparison of volatile organic emissions at the beginning and end of evaporator batches.

The top 20 SVOCs measured in the offgas stream are plotted in Figure 8. Semivolatile organic compounds also appear to be only slightly higher at the start of the evaporator batch. Benzoic acid (a target analyte) and benzaldehyde (a tentatively identified compound) were the two most prevalent semivolatile organics emitted during operation of the NWCF ETS. The maximum emissions concentrations for benzoic acid and benzaldehyde were 310 ppb and 80 ppb, respectively. On a volumetric basis, the sum of all volatile and semivolatile organics is less than 1 ppm. All other SVOCs measurements are near the method detection limits for the respective species.

With the exception of benzoic acid, all of the SVOC species emitted from the evaporator were also detected during the NWCF Calciner offgas emissions inventory (Boardman 2001). Nearly all of the compounds are derivatives of benzene or other cyclic compounds and are possibly the products of incomplete combustion of the kerosene used to heat the Calciner. These compounds probably entered the tank system when Calciner scrub was recycled to the tank farm. It is further postulated that benzoic acid and benaldehyde were either formed during combustion of the kerosene or they were formed by oxidation of benzene and toluene in the acidic waste solutions. Relatively higher emissions of benzoic acid can be explained by noting that it is readily stripped from waste solutions by steam.

In summary, the rate of organic emissions at the start and end of the evaporator batches were not significantly different. This phenomena is attributed to the trade off between evaporator temperature and the volume of fresh waste solution fed to the evaporator during the respective sampling periods. The hourly total emissions rate for all volatile and semivolatile organic emissions was less than 0.02 lbs/hr. This is significantly less than the 3 lbs/hr limit that is generally considered significant for RCRA waste treatment units permitting decisions.

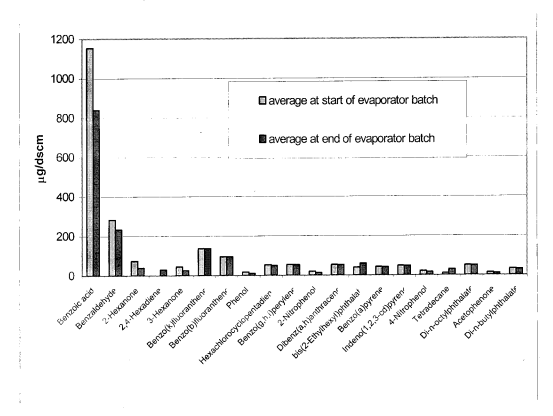


Figure 8. Comparison of semi-volatile organic emissions at the beginning and end of evaporator batches

# 9.1.2 Inorganic Compounds

The average metals emissions rates at the beginning and end of evaporator batches are plotted in Figure 9. As anticipated, metals emissions, including mercury, were typically higher at the end of an evaporator batch when the evaporator solution reached its maximum density. Assuming aerosol droplet entrainment with the overhead gas was constant throughout the run, then the emissions of all nonvolatile species should correlate with the solution density. The exception is mercury. If volatile elemental mercury exists as a dissolved gas in the waste solutions, then it would tend to be volatilized at the beginning of the evaporator runs. Otherwise, if the mercury is complexed or speciated in the wastes, then it would tend to be emitted as a non-volatile entrained species. These data indicate that mercury was mainly emitted as non-volatile particulate, although no speciation was attempted to distinguish elemental versus oxidized forms of mercury in the effluent gas stream. Aluminum, manganese, and zinc emissions appear to correlate with their relative abundance in the evaporator feed and bottoms. The emissions of all other metals were relatively low, as were their concentrations in the feed.

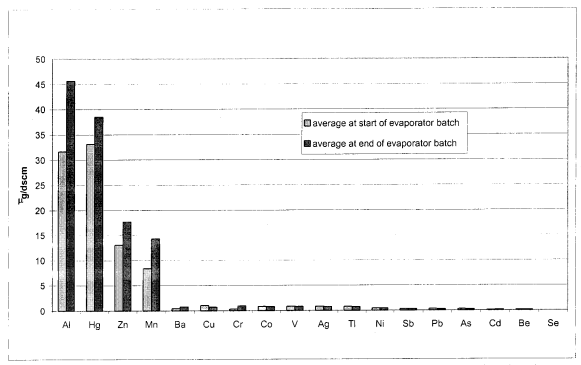


Figure 9. Comparison of metals emissions at the beginning and end of evaporator batches (not including final probe rinse species apportionment).

Throughout the sampling period, it was necessary to leave the 12-ft Method 5 sample collection probe at a fixed position in the offgas duct to reduce the potential for contamination spread and personnel radiation exposures. The probe was removed at the end of the 3-week sampling period and rinsed with acetone, followed by nitric acid, to obtain a final probe rinse measurement of particulate and metals absorbed on the probe's glass liner. Approximately 93 dscm of offgas sample was drawn through the probe over the duration of the offgas emissions inventory. The apportioned amount of particulate and metals for a single run is thus roughly 3/93 [dscm/dscm] or 3.2% of the total particulate and metals mass measured in the final probe rinse. Thus, 3.2% of the particulate and metals detected in the final probe rinses should be apportioned to the run averages.

The difference between the four most abundant metals detected in the offgas and the apportioned amount in the final probe rinse, relative to the average emissions was <10% for Zn, <6% for Al, <0.5% for Mn, and <0.1% for Hg. The percent of apportioned probe mass for the minor species was also typically low, although a comparison of the results is skewed by the fact that the measurements are near or below the analytical method detection limits. In conclusion, these results indicate that the metals uptaken on the probe liner were insignificant with respect to obtaining an accurate emissions inventory for the target metals species.

Total particulate and chloride emissions rate averages at the start and end two evaporator batches are illustrated in Figures 10 and 11. There were relatively small differences in the chloride emissions at the start and end of the batch. Hydrochloric acid levels are significantly higher as expected. Still, the sum of chloride emission contributions from HCl and Cl<sub>2</sub> is less than 1 ppmv.

Particulate emissions follow the trend of the semi-volatile organic species which were slightly higher at the beginning of the batch. Figure 11 also compares the apportioned particulate measurement for the final probe rinse (*viz.*, 3.2% of the final probe particulate measurement as discussed above). The relative amount of particulate absorbed on the probe was 20-25% of the train total. It can be inferred that some SVOCs were also deposited on the probe liner, with the maximum being similar to the particulate. This fact should be taken into consideration when using the SVOC emissions rates.

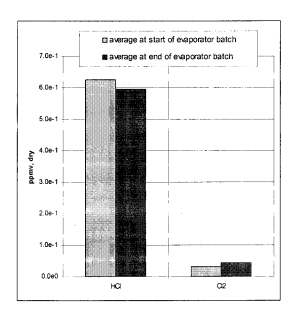


Figure 10. Comparison of chloride emissions at the beginning and end of evaporator batches.

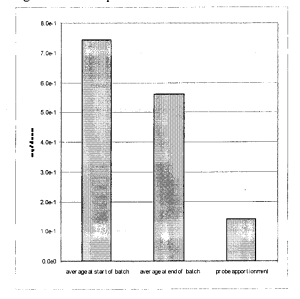


Figure 11. Comparison of particulate emissions at the beginning and end of the evaporator run with probe particulate apportionment.

## 9.2 Emissions Health Risk

The emission rates of hazardous air pollutants from the NWCF ETS were used to calculate risk to human health. Pollutants from the NWCF ETS are released from the same point and under the same conditions as NWCF Calciner emissions. Therefore, to a close approximation, the NWCF ETS can be scaled to the risk terms previously determined for the NWCF Calciner. The NWCF Calciner results are described in NWCF Calciner Emissions Inventory -Final Report for Phase IV Testing (2001).

Compounds with an EPA hazards quotient (HQ) or a cancer risk (Risk) present in the NWCF ETS samples, were ratioed to the NWCF Calciner emissions rate to determine an estimated NWCF ETS risk. Maximum values were used to bound risks. The NWCF Calciner emission rates were normalized to an annual basis. To compare the NWCF ETS results, measured NWCF ETS emissions were multiplied by a factor of 0.274 to normalize them to an annual basis. This is based on the NWCF ETS operating twelve hours a day, 200 days a year.

It was observed that the emissions rates were much lower for the NWCF ETS than from the NWCF Calciner with the exception of benzoic acid. The semi-volatiles were the largest contributor to the HQ and the Risk. The largest contributor was a phthalate (bis(2 ethylhexyl)phthalate) which is a common sampling or laboratory contaminate from plastics such as tubing, bottles, etc. Most of the materials "detected" were present at levels below the RL and in only a few samples. Benzoic acid, the single organic found in high concentrations than the NWCF Calciner, has a relatively low cancer risk and hazard quotient compound. The total HQ for the NWCF ETS was 6.2e-6 as compared to 3.3 e-03 for the NWCF Calciner and 0.25 for the EPA target criteria. The cancer risk was 1.3e-10 compared to 1.9e-07 for the NWCF Calciner and an EPA target of 1e-5.

A risk summary is given in Figures 12 and 13. Tables 18, 19, 20, and 21 give the concentration ratios and scaled risk factors.

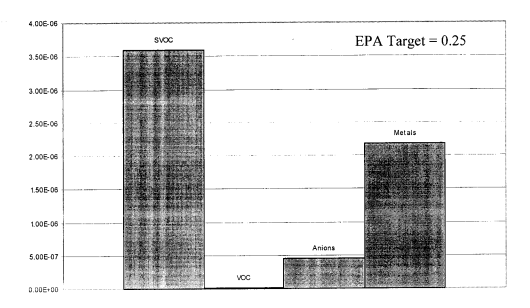


Figure 12. NWCF ETS EPA hazards quotient.

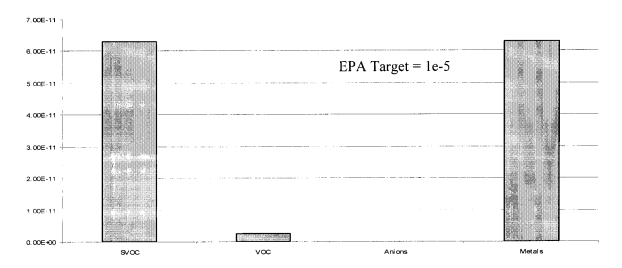


Figure 13. NWCF ETS cancer risk by pollutant category.

Table 18. Risk scaling of Method 0010 analytes.

Analyte	NWCF Hazard	Concentration Ratio	NWCF ETS Hazard	Risk
Acenaphthene	3.70E-12	2.61	9.60E-12	
Acenaphthylene				
Acetophenone	5.60E-09	0.0481	2.69E-10	
Anthracene	6.60E-12	351	2.31E-10	
Benzoic acid	4.50E-09	0.15	6.75E-10	
Benzo(a)anthracene				
Benzo(a)pyrene				
Benzo(b)fluoranthene				
Benzo(g,h,i)perylene				
Benzo(k)fluoranthene				
bis(2-Chloroethyl)ether	9.90E-10	0.02	1.98E-11	2.00E-11
bis(2-Ethylhexyl)phthalate	1.70E-04	0.013	2.21E-06	2.00E-11
4-Bromophenyl-phenylether	4.60E-07	0.03	1.38E-08	
Butylbenzylphthalate	7.50E-09	0.02	1.50E-10	
Carbazole				
2-Chloronaphthalene	4.20E-09	0.02	8.40E-11	
2-Chlorophenol	2.20E-06	8.00E-04	1.76E-09	
4-Chlorophenyl phenyl ether				
Chrysene				
Di-n-butylphthalate	4.50E-07	0.025	1.13E-08	
Di-n-octylphthalate	3.50E-09	0.1	3.50E-10	
Dibenz(a,h)anthracene				
Dibenzofuran				
1,2-Dichlorobenzene	1.60E-09	0.09	1.44E-10	
1,3-Dichlorobenzene	9.90E-10	0.1	9.90E-11	
1,4-Dichlorobenzene	3.90E-10	0.12	4.68E-11	
Diethylphthalate	3.20E-09	0.02	6.40E-11	
Dimethyl phthalate	1.40E-10	2.00E-02	2.80E-12	
2,4-Dinitrotoluene	3.90E-08	0.014	5.46E-10	
2,6-Dinitrotoluene	2.10E-06	0.02	4.20E-08	
1,2-Diphenylhydrazine	3.50E-09	0.02	7.00E-11	8.00E-12
Fluoranthene	9.70E-10	21.51	2.09E-08	
Fluorene				
Hexachlorocyclopentadiene	2.40E-04	2.00E-03	4.80E-07	
Hexachlorobenzene	6.10E-07	0.02	1.22E-08	1.50E-1
Hexachlorobutadiene				
Hexachloroethane	8.20E-07	0.014	1.15E-08	1.50E-1
Indeno(1,2,3-cd)pyrene				
Isophorone	3.90E-09	0.02	7.80E-11	1.00E-14
2-Methylnaphthalene				

Table 18. Risk scaling of Method 0010 analytes.

Analyte	NWCF	Concentration	NWCF ETS	Risk
	Hazard	Ratio	Hazard	
N-Nitrosodimethylamine				
N-Nitrosodiphenylamine	2.40E-12	0.03	7.20E-14	
Naphthalene	3.60E-07	0.014	5.04E-09	
Nitrobenzene	3.40E-09	3.00E-03	1.02E-11	
2-Nitrophenol				
4-Nitrophenol	1.40E-06	8.00E-04	1.12E-09	
2,2'-Oxybis(1-chloropropane)				
Phenanthrene				
Phenol	2.90E-08	4.00E-03	1.16E-10	
Pyrene	3.20E-08	341	1.09E-06	
1,2,4-Trichlorobenzene	1.60E-09	0.1	1.60E-10	
Total		-	3.90E-06	6.32E-11

1. Compounds evaluated using PAH high resolution method for NWCF Calciner.

Table 19. Risk scaling for Method 0031 analytes.

Analyte	NWCF	Concentration	NWCF ETS	Risk
	Hazard	Ratio	Hazard	
Acetone	6.30E-08	0.023	1.45E-09	
Benzene	4.10E-06	7.70E-04	3.16E-09	
Bromomethane				
2-Butanone	1.50E-09	0.046	6.90E-11	
Carbon disulfide	9.00E-10	0.147	1.32E-10	
Carbon tetrachloride	1.30E-08	0.0351	4.56E-10	1.70E-13
Chlorobenzene	4.80E-08	0.0076	3.65E-10	
Chloroethane	3.20E-11	0.044	1.41E-12	
Chloroform	1.90E-08	0.044	8.36E-10	6.60E-13
Chloromethane	1.50E-09	0.466	6.99E-10	3.70E-13
Dichlorodifluoromethane	2.00E-09	0.0466	9.32E-11	
1,2-Dichloroethane	3.20E-08	0.0356	1.14E-09	3.00E-13
1,1-Dichloroethene	1.00E-08	0.041	4.10E-10	3.40E-13
1,2-Dichloropropane	8.10E-08	0.03	2.43E-09	1.90E-13
Methylene chloride	5.50E-10	0.877	4.82E-10	4.80E-13
Toluene	7.20E-10	0.055	3.96E-11	
Trichlorofluoromethane	8.30E-10	0.02	1.66E-11	
Vinyl chloride	1.80E-12	0.0356	6.41E-14	6.40E-14
o-Xylene	4.80E-11	0.018	8.64E-13	
Totals			1.18E-08	2.574E-12

Table 20. Risk scaling for Method 0050 analytes.

Analyte	NWCF Hazard	Concentration	NWCF ETS	Risk
		Ratio	Hazard	
***************************************				
Chloride (as HCl)	1.50E-05	0.02	3.00E-07	
Chloride (as Cl2)	1.70E-07	0.0047	7.99E-10	
Fluoride (as HF)	3.50E-05	0.00082	2.87E-08	
Nitrate (as HNO3)	9.50E-04	0.00008	7.60E-08	
Nitrite (as HNO2)				
Particulate				
Total			4.05E-07	

Table 21. Risk scaling for Method 0060 analytes.

Analyte	NWCF Hazard	Concentration	HLLWE Hazard	Risk
		Ratio		
Aluminum (Al)	5.7E-09	0.036	2.1E-10	
Antimony (Sb)	1.50E-09	0.056	8.48E-11	
Arsenic (As)	1.30E-14	2.2E-7	7.8E-22	
Barium (Ba)	7.50E-06	0.0025	1.88E-08	
Beryllium (Be)				
Cadmium (Cd)	1.70E-07	0.026	4.5E-09	1.70E-12
Chromium (Cr)	9.00E-07	0.047	4.23E-08	6.0E-11
Cobalt (Co)				
Copper (Cu)	4.50E-09	0.017	7.5E-11	
Lead (Pb)	2.90E-08	0.008	2.4E-10	
Manganese (Mn)	1.60E-05	0.085	1.36E-06	
Mercury (Hg)	2.10E-04	0.0036	7.56E-07	
Nickel (Ni)	2.30E-11	0.07	1.61E-12	1.30E-12
Selenium (Se)	8.20E-10	2.00E-11	1.64E-20	
Silver (Ag)				
Thallium (Tl)	2.7E-8	0.14	4.0E-9	
Vanadium (V)				
Zinc (Zn)	2.00E-08	5.00E-05	1.00E-12	
		•	Total	
Total			2.19E-06	6.3E-11

### 10. CONCLUSIONS

Characterization samples for the NWCF ETS were collected with only minor deviations from EPA protocols. Due to ALARA concerns, the samples were collected at a single point in the duct and the probe was not removed between sample trains. The NWCF ETS emissions rates for all species were relatively low in terms of regulatory emissions limits and health risk considerations. It was observed that organic compound emissions are slightly higher at the beginning of the batch while metals emissions, including mercury, are slightly higher at the end of the evaporator batch. Mercury emissions were less than 5 ppbv (< 40  $\mu$ g/dscm), while the sum of HCl and Cl2 emissions was less than 1 ppmv. The sum of all organic emissions also was less than 1 ppmv. Particulate emissions (included the apportioned particulate recovered in the final probe rinse) are less than 0.9 mg/dscm and less than 0.7 mg/dscm at the beginning and end of the evaporator batch, respectively.

The estimated HQ for the evaporator was 6.2e-6 as compared to 0.25 for the EPA target criteria. The estimated cancer risk was 1.3e-10 compared to an EPA target of 1.0e-5. The NWCF ETS offgas emissions inventory was completed in accordance with the QAPjP developed and approved for this activity. Conventional EPA sampling and analytical methods were used to characterize volatile and semivolatile organic compounds, multiple metals, HCl/Cl<sub>2</sub>, and particulate emissions.

Diligence in following sample checklists, continuous monitoring by either the Project Technical Leads and Quality Assurance Office, use of a master sample collection list, pre-defined sample labels, and RFA/COC documentation provided for the best possible sample collection accuracy and consistency. The data are believed to be accurate and representative of the NWCF ETS for the feed and system operating conditions during the offgas sampling period. A compilation of the process operating parameters, the offgas sample analytical data summaries, and the calculated emissions rates and liquid composition data for the evaporator feed, overhead condensate, and bottoms are included in the report appendices for permit applications purposes.

NWCF ETS operations were normal and consistent throughout the three-week sample collection period. Feed batches were consistent, as were the evaporator operating parameters and offgas system conditions. Radiation levels in the offgas samples were extremely low. Extended gamma scanning did not identify any gamma emitters in either the sample contamination trains or ongoing screening samples. Gross beta and gross alpha levels were only detected in the pico-curies range, easily meeting all of the analytical laboratory sample acceptance criteria. Tritium levels were low and proportionate to the low levels of moisture that were present in the NWCF offgas stream. Oxygen levels in the offgas duct were comparable to ambient air conditions.

Species absorption in the probe liner was minor. The exception may be organic particulate. The apportioned amount of organic particulate contained in the final probe rinse was approximately 25% of the average Method 0050 run total particulate measurements. This suggests that some semi-volatile organic matter could be potentially deposited on the probe liner. Even when the SVOC results are conservatively escalated by 25%, to account for the maximum potential portion of semivolatile material adsorbed on the probe liner, the outcome of the emissions rates are risk calculations are not significant.

Metal adsorption on the probe was low for all metals. Less than 0.1% of the mercury was deposited on the probe liner. Therefore, apportionment of the final probe rinse to the Method 0060 Metals trains is not significant.

#### 11. REFERENCES

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BECHTEL BWXT IDAHO, LLC (BBWI)
INTEC HLLWE Effluent Gas Emissions Inventory
Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### MM-5 Train Summary - Run 1 Train Totals Semivolatile Organic Compounds Analytical Results Summary Table A-1. HLLWE Run ID: 0010-STRT-1

Field Sample Name:

MM-5 Train

Sample Description:

MM-5 Train Totals for Semivolatile Organic Compounds Analysis

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (µg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (µg)		MM-5 Tota (Tota	Project Specific	
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag	Risk Result	Flag	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
Target Compound List			,							
Acenaphthene	83-32-9	1.3	U	6.6	U	1.9	U	< 9.8		N
Acenaphthylene	208-96-8	1.3	U	6.6	U	1.6	U	< 9.5		N
Acetophenone	9[8-86-2	2.0	U	44	J	2.6	U	< 49	J	Р
Aniline	62-53-3	2.5	U	94	U	18	U	< 110		N
Anthracene	120-12-7	1.3	U	6.6	U	1.6	U	< 9.5		N
Benzidine	92-87-5	100	U	500	U	66	U	< 670		N
Benzoic acid	65-85-0	100	U	4,800	Е	9.4	U	< 4,900	Е	P
Benzo(a)anthracene	56-55-3	2.2	U	7.6	U	1.7	U	< 12		N
Benzo(a)pyrene	50-32-8	2.6	U	130	U	1.8	U	< 130		N
Benzo(b)fluoranthene	205-99-2	3.7	U	290	U	4.2	U	< 300		N
Benzo(g,h,i)perylene	191-24-2	7.3	U	160	U	2.1	U	< 170		N
Benzo(k)fluoranthene	207-08-9	5.5	U	420	U	2.9	U	< 430		N
Benzyl alcohol	100-51-6	92	U	470	U	4.2	U	< 570		N
bis(2-Chloroethoxy)methane	111-91-1	1.5	U	6.6	U	2.0	U	< 10		N
bis(2-Chloroethyl)ether	111-44-4	2.0	U	7.3	U	1.7	U	< 11		N
bis(2-Ethylhexyl)phthalate	117-81-7	14		100	J	11		< 120	J	A
4-Bromophenyl-phenylether	101-55-3	1.4	U	6.6	U	1.4	U	< 9.4		N
Butylbenzylphthalate	85-68-7	2.9	U	7.9	U	2.3	U	< 13		N
Carbazole	86-74-8	2.0	U	8.4	U	2.2	U	< 13		N
4-Chloro-3-methylphenol	59-50-7	2.6	U	8.1	U	6.6	U	< 17		N
4-Chloroaniline	106-47-8	3.1	U	79	U	7.9	U	< 90		N
2-Chloronaphthalene	91-58-7	1.3	U	6.6	U	1.4	U	< 9.3		N
2-Chlorophenol	95-57-8	2.6	U	6.6	U	1.7	U	< 11		N
4-Chlorophenyl phenyl ether	7005-72-36	1.3	U	6.6	U	3.1	U	< 11		N
Chrysene	218-01-9	2.3	U	8.4	U	1.3	U	< 12		N

## MM-5 Train Summary - Run 1 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-1. HLLWE Run ID: 0010-STRT-1

	CAS Registry	Front H Composi	MM-5 Train Front Half Composite <sup>1</sup> (μg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)	
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
Di-n-butylphthalate	84-74-2	1.9	J	100	U	2.3	U	< 100	J	P
Di-n-octylphthalate	117-84-0	11		150	U	2.5	J	< 160	J	P
Dibenz(a,h)anthracene	53-70-3	5.2	U	160	U	2.9	U	< 170		N
Dibenzofuran	132-64-9	1.4	U	6.6	U	2.9	U	< 11		N
1,2-Dichlorobenzene	95-50-1	2.2	U	6.8	U	1.7	U	< 11		N
1,3-Dichlorobenzene	541-73-1	3.1	U	7.3	U	1.4	U	< 12		N
1,4-Dichlorobenzene	106-46-7	2.9	U	11	J	2.0	U	< 16	J	Р
3,3'-Dichlorobenzidine	91-94-1	7.1	U	97	U	7.9	U	< 110		N
2,4-Dichlorophenol	120-83-2	3.9	U	6.6	U	2.3	U	< 13		N
Diethylphthalate	84-66-2	5.4	J	9.4	U	1.4	U	< 16	J	Р
Dimethyl phthalate	131-11-3	1.7	U	6.6	U	1.3	U	< 9.6	•	N
2,4-Dimethylphenol	105-67-9	7.6	U	50	U	1.5	U	< 59		N
4,6-Dinitro-2-methylphenol	534-52-1	13	U	120	U	1.5	U	< 130		N
2,4-Dinitrophenol	51-28-5	15	U	250	U	3.9	U	< 270		N
2,4-Dinitrotoluene	121-14-2	4.2	U	6.6	U	2.6	U	< 13		N
2.6-Dinitrotoluene	606-20-2	3.4	U	6.6	U	2.1	U	< 12		N
1,2-Diphenylhydrazine	122-66-7	1.7	U	6.6	U	1.5	U	< 9.8		N
Fluoranthene	206-44-0	1.3	U	7.1	U	1.8	U	< 10		N
Fluorene	86-73-7	1.3	U	6.6	U	2.6	U	< 10		N
Hexachlorocyclopentadiene	77-47-4	26	U	130	U	6.6	U	< 160		N
Hexachlorobenzene	118-74-1	1.5	U	6.6	U	2.6	U	< 11		N
Hexachlorobutadiene	87-68-3	3.7	U	9.7	U	1.9	U	< 15		N
Hexachloroethane	67-72-1	6.6	U	7.1	U	1.9	U	< 16		N
Indeno(1,2,3-cd)pyrene	193-39-5	5.5	U	140	U	2.3	U	< 150		N
Isophorone	78-59-1	1.7	U	6.6	U	1.8	U	< 10		N
2-Methylnaphthalene	91-57-6	1.5	U	6.6	U	2.3	U	< 10		N
2-Methylphenol	95-48-7	6.0	U	39	U	2.1	U	< 47		N
3-Methylphenol & 4-Methylphenol	65794-96-9	6.0	U	26	U	2.2	U	< 34		N

## MM-5 Train Summary - Run 1 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-1. HLLWE Run ID: 0010-STRT-1

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (μg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>		Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
N-Nitroso-di-n-propylamine	621-64-7	1.9	IJ	6.6	U	2.3	U	< 11		N
N-Nitrosodimethylamine	62-75-9	1.9	U	6.6	U	2.2	U	< 11		N
N-Nitrosodiphenylamine	86-30-6	1.6	U	12	U	1.4	U	< 15		N
Naphthalene	91-20-3	1.3	U	7.9	U	1.8	U	< 11		N
2-Nitroaniline	88-74-4	1.5	U	6.6	U	2.9	U	< 11		N
3-Nitroaniline	99-09-2	10	U	26	U	4.7	U	< 41		N
4-Nitroaniline	100-01-6	6.0	U	26	U	3.9	U	< 36		N
Nitrobenzene	98-95-1	1.9	U	12	J	1.7	U	< 16	J	P
2-Nitrophenol	88-75-5	8.4	U	80		2.6	U	< 91		P
4-Nitrophenol	100-02-7	8.7	U	63	J	3.9	U	< 76	J	Р
2,2'-Oxybis(1-chloropropane) 8	108-60-1	2.6	U	10	U	1.8	U	< 14		N
Pentachlorobenzene	608-93-5	1.4	U	6.6	U	2.3	U	< 10		N
Pentachloronitrobenzene	82-68-8	2.0	U	6.6	U	2.6	U	< 11		N
Pentachlorophenol	87-86-5	50	U	250	U	3.4	U	< 300		N
Phenanthrene	85-01-8	1.3	U	6.6	U	1.9	· U	< 9.8		N
Phenol	108-95-2	2.9	U	72		2.2	U	< 77		P
Pyrene	129-00-0	1.9	U	6.8	U	1.4	U	< 10		N
Pyridine	110-86-1	2.3	U	9.7	U	5.2	U	< 17		N
1,2,4,5-Tetrachlorobenzene	95-94-3	2.3	U.	6.6	U	2.2	U	< 11		N
1,2,4-Trichlorobenzene	120-82-1	1.9	U	7.9	U	2.2	U	< 12		N
2,4,5-Trichlorophenol	95-95-4	6.0	U	17	U	2.1	U	< 25		N
2,4,6-Trichlorophenol	88-06-2	3.7	U	10	U	2.5	U	< 16		N

## MM-5 Train Summary - Run 1 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-1. HLLWE Run ID: 0010-STRT-1

CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (ug)	MM-5 Train Back Half Composite <sup>2</sup> (µg)	MM-5 Train Condensate Composite <sup>3</sup> (μg)	MM-5 Train Totals <sup>4</sup> (Total µg)		Project Specific
Number				Total <sup>6</sup>	Flag	Flag <sup>7</sup>
625-86-5	4.1		9.5	14	N,J,M	Р
589-38-8		190		190	N,J,M	P
591-78-6		230		230	N,J,M	P
2216-33-3	7.9			7.9	N,J,M	P
100-52-7		1,100		1,100	N,J,M	P
112-40-3		67		67	N,J,M	Р
629-50-5		20		20	N,J,M	Р
629-59-4	8.6			8.6	N,J,M	P
629-62-9	15			15	N,J,M	Р
126-73-8	23			23	N,J,M	P
294-62-2	20		11	31	N,J,M	P
629-78-7	5.5		3.4	8.9	N,J,M	P
112-95-8			2.0	2.0	N,J,M	P
57-10-3	5.4			5.4	N,J,M	Р
57-11-4	3.6			3.6	N,J,M	Р
791-28-6			8.7	8.7	N,J,M	Р
629-94-7			2.0	2.0	N,J,M	P
7098-22-8			6.6	6.6	N,J,M	P
112-95-8			4.1	4.1	N,J,M	Р
	Registry Number  625-86-5 589-38-8 591-78-6 2216-33-3 100-52-7 112-40-3 629-50-5 629-59-4 629-62-9 126-73-8 294-62-2 629-78-7 112-95-8 57-10-3 57-11-4 791-28-6 629-94-7 7098-22-8	CAS Registry Number  625-86-5  625-86-5  591-78-6  2216-33-3  7.9  100-52-7  112-40-3  629-50-5  629-59-4  8.6  629-62-9  15  126-73-8  23  294-62-2  20  629-78-7  57-10-3  57-11-4  3.6  791-28-6   629-94-7  7098-22-8	CAS Registry Number         Front Half Composite¹ (μg)         Back Half Composite² (μg)           625-86-5         4.1            589-38-8          190           591-78-6          230           2216-33-3         7.9            100-52-7          1,100           112-40-3          67           629-50-5          20           629-59-4         8.6            629-62-9         15            126-73-8         23            294-62-2         20            57-10-3         5.4            57-11-4         3.6            791-28-6             629-94-7             7098-22-8	CAS Registry Number         Front Half Composite¹ (μg)         Back Half Composite² (μg)         Condensate Composite³ (μg)           625-86-5         4.1          9.5           589-38-8          190            591-78-6          230            100-52-7          1,100            629-50-5          20            629-59-4         8.6             629-62-9         15             126-73-8         23             294-62-2         20          11           629-78-7         5.5          3.4           112-95-8           2.0           57-10-3         5.4             57-11-4         3.6             791-28-6           8.7           629-94-7           2.0	CAS Registry Number         Front Half Composite (μg)         Back Half Composite (μg)         Condensate Composite (μg)         MM-5 Total (Total (Tota	CAS Registry Number         Front Half Composite <sup>1</sup> (μg)         Back Half Composite <sup>2</sup> (μg)         Condensate Composite <sup>3</sup> (μg)         MM-5 Train Totals <sup>4</sup> (Total μg)           625-86-5 89-38-8         4.1          9.5         14         N,J,M           591-78-6          230          230         N,J,M           100-52-7          1,100          1,100         N,J,M           629-50-5          20          20         N,J,M           629-62-9         15          15         N,J,M           629-78-7         5.5          23         N,J,M           629-78-7         3.4         8.9         N,J,M           112-95-8           2.0         2.0         N,J,M           57-10-3         5.4           5.4         N,J,M           791-28-6           2.0         2.0         N,J,M           7098-22-8           8.6         N,J,M           7098-22-8           2.0         2.0         N,J,M           7098-22-8

#### Footnotes:

- The MM-5 Train Front Half Composite consists of the Particulate Filter and the Front Half of the Filter Holder and Probe Solvent Rinses.
- The MM-5 Train Back Half Composite consists of the XAD-2 Resin Tube and the Back Half of the Filter Holder and Coil Condenser Solvent Rinses.
- <sup>3</sup> The MM-5 Train Condensate Composite consists of the Condensate and Impinger Contents and the Glassware Solvent Rinses.
- The total mass for each semivolatile compound found in the MM-5 sampling train consists of the sum of the MM-5 train's Front Half Composite contents, the train's Back Half Composite contents, and the Condensate Composite. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) + (Concentration in the Condensate Composite x Condensate Composite Volume) = Total  $\mu g$  in the MM-5 Sampling Train.

Therefore:  $(\mu g) + (\mu g) + (\mu g/Liter \times Liter) = Total \mu g$ 

The MM-5 Train Run Total (in Total  $\mu g$ ) is the sum of results for the three (3) MM-5 train sample fractions using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
- ♦ When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MM-5 Train Total are the cumulative set of flags for each train component included as part of the MM-5 train total. A flag attached to an MM-5 train component is carried through to the "MM-5 Train Total" column when the associated component analytical result is a significant number in comparison to the MM-5 Train Total. That is, if the MM-5 Train Total is affected by an MM-5 train component analytical result, the flag is carried through to the MM-5 Train Total, but if the MM-5 Train Total is not affected by an MM-5 train component, the flag is not carried through to the MM-5 Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "J" qualifier indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - ♦ A "B" qualifier indicates that this compound was found in the associated laboratory method blank. Under these conditions these values are regarded as estimated values.
  - A "D" qualifier indicates that this result was obtained through dilution of the sample. This original analysis yielded a result that exceeded the calibration range.
  - An "N" qualifier indicates that this compound is a tentatively identified compound (TIC). Therefore the value is estimated.
  - An "E" qualifier indicates that this compound exceeded the calibration range of the instrument.
  - An "A" qualifier indicates that this result is an Aldol-condensation product.
  - An "M" qualifier indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- <sup>8</sup> Bis(2-chloroisopropyl)ether and 2,2'-Oxybis(1-chloropropane) are synonyms.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data. It should be noted that TICs that give the same mass spectral match for GC peaks at different retention times are listed separately with the same compound identity. Under these conditions the compounds are likely indistinguishable isomers of the same compound. However, insufficient evidence is available to determine unequivocal identities.

#### MM-5 Train Summary - Run 3 Train Totals Semivolatile Organic Compounds Analytical Results Summary Table A-2. HLLWE Run ID: 0010-END-1

Field Sample Name:

MM-5 Train

Sample Description:

MM-5 Train Totals for Semivolatile Organic Compounds Analysis

	CAS Registry	MM-5 Ti Front H Compos (µg)	alf	MM-5 Tr Back Ha Composi (μg)	alf ite <sup>2</sup>	MM-5 Ti Condens Compos (μg)	ate ite <sup>3</sup>	MM-5 Tota (Tota	als <sup>4</sup> I μg)	Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
Target Compound List										
Acenaphthene	83-32-9	1.3	U	6.6	U	1.9	U	< 9.8		N
Acenaphthylene	208-96-8	1.3	U	6.6	U	1.6	U	< 9.5		N
Acetophenone	9[8-86-2	2.0	U	31	J	2.6	U	< 36	J	P
Aniline	62-53-3	2.5	U	94	U	18	U	< 110		N
Anthracene	120-12-7	1.3	U	6.6	U	1.6	U	< 9.5		N
Benzidine	92-87-5	100	U	500	U	66	U	< 670		N
Benzoic acid	65-85-0	100	U	2,500	Е	9.4	U	< 2,600	Е	P
Benzo(a)anthracene	56-55-3	2.2	U	7.6	U	1.7	U	< 12		N
Benzo(a)pyrene	50-32-8	2.6	U	130	U	1.8	U	< 130		N
Benzo(b)fluoranthene	205-99-2	3.7	U	290	U	4.2	U	< 300		N
Benzo(g,h,i)perylene	191-24-2	7.3	U	160	U	2.1	U	< 170		N
Benzo(k)fluoranthene	207-08-9	5.5	U	420	U	2.9	U	< 430		N
Benzyl alcohol	100-51-6	92	U	470	U	4.2	U	< 570		N
bis(2-Chloroethoxy)methane	111-91-1	1.5	U	6.6	U	2.0	U	< 10		N
bis(2-Chloroethyl)ether	111-44-4	2.0	U	7.3	U	1.7	U	< 11		N
bis(2-Ethylhexyl)phthalate	117-81-7	49		100	U	68	,,	< 220		P
4-Bromophenyl-phenylether	101-55-3	1.4	U	6.6	U	1.4	U	< 9.4		N
Butylbenzylphthalate	85-68-7	2.9	U	7.9	U	2.3	U	< 13		N
Carbazole	86-74-8	2.0	U	8.4	U	2.2	U	< 13		N
4-Chloro-3-methylphenol	59-50-7	2.6	U	8.1	U	6.6	U	< 17		N
4-Chloroaniline	106-47-8	3.1	U	79	U	7.9	U	< 90		N
2-Chloronaphthalene	91-58-7	1.3	U	6.6	U	1.4	U	< 9.3		N
2-Chlorophenol	95 <b>-</b> 57-8	2.6	U	6.6	U	1.7	U	< 11		N
4-Chlorophenyl phenyl ether	7005-72-36	1.3	U	6.6	U	3.1	U	< 11		N
Chrysene	218-01-9	2.3	U	8.4	U	1.3	U	< 12		N

## MM-5 Train Summary - Run 3 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-2. HLLWE Run ID: 0010-END-1

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (μg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
	04.74.3	1.0	T	100	U	2.3	U	< 100	J	P
Di-n-butylphthalate	84-74-2	1.9	J J	150	U	2.5	J	< 160	J	P
Di-n-octylphthalate	117-84-0	8.4			U	2.9	U	< 170	J	N
Dibenz(a,h)anthracene	53-70-3	5.2	U	160	U	2.9	U	< 11		N
Dibenzofuran	132-64-9	1.4	U	6.6				< 11		N
1,2-Dichlorobenzene	95-50-1	2.2	U	6.8	U	1.7	U	< 12		N
1,3-Dichlorobenzene	541-73-1	3.1	U	7.3	U	1.4	U		J	P
1,4-Dichlorobenzene	106-46-7	2.9	U	6.8	J	2.0	U	< 12	J	N
3,3'-Dichlorobenzidine	91-94-1	7.1	U	97	U	7.9	U	< 110		
2,4-Dichlorophenol	120-83-2	3.9	U	6.6	U	2.3	U	< 13		N
Diethylphthalate	84-66-2	3.9	U	9.4	U	1.4	U	< 15	<u> </u>	N
Dimethyl phthalate	131-11-3	1.7	U	6.6	U	1.3	U	< 9.6		N
2,4-Dimethylphenol	105-67-9	7.6	U	50	U	1.5	U	< 59	<u> </u>	N
4,6-Dinitro-2-methylphenol	534-52-1	13	U	120	U	1.5	U	< 130		N
2,4-Dinitrophenol	51-28-5	16	U	250	U	3.9	U	< 270		N
2,4-Dinitrotoluene	121-14-2	4.2	U	6.6	U	2.6	U	< 13		N
2,6-Dinitrotoluene	606-20-2	3.4	U	6.6	U	2.1	U	< 12		N
1,2-Diphenylhydrazine	122-66-7	1.7	U	6.6	U	1.5	U	< 9.8		N
Fluoranthene	206-44-0	1.3	U	7.1	U	1.8	U	< 10		N
Fluorene	86-73-7	1.3	U	6.6	U	2.6	U	< 10		N
Hexachlorocyclopentadiene	77-47-4	26	U	130	U	6.6	U	< 160		N
Hexachlorobenzene	118-74-1	1.5	U	6.6	U	2.6	U	< 11		N
Hexachlorobutadiene	87-68-3	3.7	U	9.7	U	1.9	U	< 15		N
Hexachloroethane	67-72-1	6.6	U	7.1	U	1.9	U	< 16		N
Indeno(1,2,3-cd)pyrene	193-39-5	5.5	U	140	U	2.3	U	< 150		N
Isophorone	78-59-1	1.7	U	6.6	U	1.8	U	< 10		N
2-Methylnaphthalene	91-57 <b>-</b> 6	1.5	U	6.6	U	2.3	U	< 10		N
2-Methylphenol	95-48-7	6.0	U	39	U	2.1	U	< 47		N
3-Methylphenol & 4-Methylphenol	65794-96-9	6.0	U	26	U	2.2	U	< 34		N

## MM-5 Train Summary - Run 3 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-2. HLLWE Run ID: 0010-END-1

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (µg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
N-Nitroso-di-n-propylamine	621-64-7	1.9	U	6.6	U	2.3	U	< 11		N
N-Nitrosodimethylamine	62-75-9	1.9	U	6.6	U	2.2	U	< 11		N
N-Nitrosodiphenylamine	86-30-6	1.6	U	12	U	1.4	U	< 15		N
Naphthalene	91-20-3	1.3	U	7.9	U	1.8	U	< 11		N
2-Nitroaniline	88-74-4	1.5	U	6.6	U	2.9	U	< 11		N
3-Nitroaniline	99-09-2	10	U	26	U	4.7	U	< 41		N
4-Nitroaniline	100-01-6	6.0	U	26	U	3.9	U	< 36		N
Nitrobenzene	98-95-1	1.9	U	11	J	1.7	U	< 15	J	P
2-Nitrophenol	88-75-5	8.4	U	30	J	2.6	U	< 41	J	Р
4-Nitrophenol	100-02-7	8.7	U	42	U	3.9	U	< 55		N
2,2'-Oxybis(1-chloropropane) 8	108-60-1	2.6	U	10	U	1.8	U	< 14		N
Pentachlorobenzene	608-93-5	1.4	U	6.6	U	2.3	U	< 10		N
Pentachloronitrobenzene	82-68-8	2.0	U	6.6	U	2.6	U	< 11		N
Pentachlorophenol	87-86-5	50	U	250	U	3.4	U	< 300		N
Phenanthrene	85-01-8	1.3	U	6.6	U	1.9	U	< 9.8		N
Phenol	108-95-2	2.9	U	32	J	2.2	J	< 37	J	P
Pyrene	129-00-0	1.9	U	6.8	U	1.4	U	< 10		N
Pyridine	110-86-1	2.3	U	9.7	U	5.2	U	< 17		N
1,2,4,5-Tetrachlorobenzene	95-94-3	2.3	U	6.6	U	2.2	U	< 11		N
1,2,4-Trichlorobenzene	120-82-1	1.9	U	7.9	U	2.2	U	< 12		N
2,4,5-Trichlorophenol	95-95-4	6.0	Ų	17	U	2.1	U	< 25		N
2,4,6-Trichlorophenol	88-06-2	3.7	U	10	U	2.5	U	< 16		N

## MM-5 Train Summary - Run 3 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-2. HLLWE Run ID: 0010-END-1

Registry	Composite <sup>1</sup> (µg)	Back Half Composite <sup>2</sup> (µg)	Condensate Composite <sup>3</sup> (µg)	MM-5 Tota (Tota	Project Specific	
Number	Risk Result Flag <sup>5</sup>			Total <sup>6</sup>	Flag	Flag <sup>7</sup>
625-86-5	4.7		3.1	7.8	N,J,M	Р
589-38-8		96		96	N,J,M	Р
591-78-6		120		120	N,J,M	Р
3074-71-3	9.9			9.9	N,J,M	Р
100-52-7		740		740	N,J,M	Р
104-57-4		72		72	N,J,M	P
4748-78-1		57		57	N,J,M	Р
112-40-3		48		48	N,J,M	P
629-50-5		21		21	N,J,M	Р
592-46-1		45		45	N,J,M	P
592-46-1		110		110	N,J,M	Р
629-59-4		96		96	N,J,M	P
126-73-8	16			16	N,J,M	Р
294-62-2	24		6.5	30	N,J,M	P
629-78-7	4.0			4.0	N,J,M	P
57-11-4	2.0			2.0	N,J,M	P
	625-86-5 589-38-8 591-78-6 3074-71-3 100-52-7 104-57-4 4748-78-1 112-40-3 629-50-5 592-46-1 592-46-1 629-59-4 126-73-8 294-62-2 629-78-7	625-86-5 4.7 589-38-8 591-78-6 3074-71-3 9.9 100-52-7 104-57-4 4748-78-1 112-40-3 592-46-1 592-46-1 629-59-4 126-73-8 16 294-62-2 24 629-78-7 4.0	625-86-5       4.7          589-38-8        96         591-78-6        120         3074-71-3       9.9          100-52-7        740         104-57-4        72         4748-78-1        57         112-40-3        48         629-50-5        21         592-46-1        45         592-46-1        96         126-73-8       16          294-62-2       24          629-78-7       4.0	625-86-5       4.7        3.1         589-38-8        96          591-78-6        120          3074-71-3       9.9           100-52-7        740          104-57-4        72          4748-78-1        57          112-40-3        48          629-50-5        21          592-46-1        45          592-46-1        96          126-73-8       16           294-62-2       24        6.5         629-78-7       4.0	625-86-5       4.7        3.1       7.8         589-38-8        96        96         591-78-6        120        120         3074-71-3       9.9         9.9         100-52-7        740        74         104-57-4        72        72         4748-78-1        57        57         112-40-3        48        48         629-50-5        21        21         592-46-1        45        45         592-46-1        96        96         126-73-8       16        96        96         126-73-78-7       4.0        6.5       30         629-78-7       4.0         4.0	625-86-5       4.7        3.1       7.8       N,J,M         589-38-8        96        96       N,J,M         591-78-6        120        120       N,J,M         3074-71-3       9.9         9.9       N,J,M         100-52-7        740        740       N,J,M         104-57-4        72        72       N,J,M         4748-78-1        57        57       N,J,M         629-50-5        21        21       N,J,M         592-46-1        45       N,J,M       110        110       N,J,M         592-46-1        96        96       N,J,M         126-73-8       16        96       N,J,M         294-62-2       24        6.5       30       N,J,M         629-78-7       4.0       N,J,M        4.0       N,J,M

#### Footnotes:

- The MM-5 Train Front Half Composite consists of the Particulate Filter and the Front Half of the Filter Holder and Probe Solvent Rinses
- <sup>2</sup> The MM-5 Train Back Half Composite consists of the XAD-2 Resin Tube and the Back Half of the Filter Holder and Coil Condenser Solvent Rinses.
- <sup>3</sup> The MM-5 Train Condensate Composite consists of the Condensate and Impinger Contents and the Glassware Solvent Rinses.
- The total mass for each semivolatile compound found in the MM-5 sampling train consists of the sum of the MM-5 train's Front Half Composite contents, the train's Back Half Composite contents, and the Condensate Composite. The calculation is as follows:

(Total µg in the Front Half) + (Total µg in the Back Half) + (Concentration in the Condensate Composite x Condensate Composite Volume)

= Total µg in the MM-5 Sampling Train.

Therefore:  $(\mu g) + (\mu g) + (\mu g/Liter \times Liter) = Total \mu g$ 

The MM-5 Train Run Total (in Total  $\mu g$ ) is the sum of results for the three (3) MM-5 train sample fractions using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
- ♦ When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MM-5 Train Total are the cumulative set of flags for each train component included as part of the MM-5 train total. A flag attached to an MM-5 train component is carried through to the "MM-5 Train Total" column when the associated component analytical result is a significant number in comparison to the MM-5 Train Total. That is, if the MM-5 Train Total is affected by an MM-5 train component analytical result, the flag is carried through to the MM-5 Train Total, but if the MM-5 Train Total is not affected by an MM-5 train component, the flag is not carried through to the MM-5 Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- <sup>5</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "J" qualifier indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" qualifier indicates that this compound was found in the associated laboratory method blank. Under these conditions these values are regarded as estimated values.
  - A "D" qualifier indicates that this result was obtained through dilution of the sample. This original analysis yielded a result that exceeded the calibration range.
  - An "N" qualifier indicates that this compound is a tentatively identified compound (TIC). Therefore the value is estimated.
  - An "E" qualifier indicates that this compound exceeded the calibration range of the instrument.
  - An "A" qualifier indicates that this result is an Aldol-condensation product.
  - An "M" qualifier indicates that this result was measured against the nearest internal standard and assumed a response factor of
    one (1).
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- <sup>8</sup> Bis(2-chloroisopropyl)ether and 2,2'-Oxybis(1-chloropropane) are synonyms.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data. It should be noted that TICs that give the same mass spectral match for GC peaks at different retention times are listed separately with the same compound identity. Under these conditions the compounds are likely indistinguishable isomers of the same compound. However, insufficient evidence is available to determine unequivocal identities.

#### MM-5 Train Summary - Run 2 Train Totals Semivolatile Organic Compounds Analytical Results Summary Table A-3. HLLWE Run ID: 0010-STRT-2

Field Sample Name:

MM-5 Train

Sample Description:

MM-5 Train Totals for Semivolatile Organic Compounds Analysis

	CAS Registry	stry (µg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 ' Tota (Total	Project Specific	
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
Target Compound List	62.20.0	1.2	U	6.6	U	8.0	J	< 16	J	P
Acenaphthene	83-32-9	1.3	U	6.6	U	7.6	J	< 16	J	P
Acenaphthylene	208-96-8	1.3		<b></b>	J	7.9	U	< 42	J	P
Acetophenone	9[8-86-2	2.0	U	32			·····	< 150	J	N
Aniline	62-53-3	2.5	U	94	U	55	U		J	P
Anthracene	120-12-7	1.3	U	6.6	U	8.3	J	< 16	J	N
Benzidine	92-87-5	100	U	500	U	200	U	< 800	-	
Benzoic acid	65-85-0	100	U	2,200	Е	29	U	< 2,300	E	P
Benzo(a)anthracene	56-55-3	2.2	U	7.6	U	9.0	J	< 19	J	P
Benzo(a)pyrene	50-32-8	2.6	U	130	U	9.3	J	< 140	J	P
Benzo(b)fluoranthene	205-99-2	3.7	U	290	U	13	J	< 300	J	P
Benzo(g,h,i)perylene	191-24-2	7.3	U	160	U	9.5	J	< 180	J	P
Benzo(k)fluoranthene	207-08-9	5.5	U	420	U	9.0	J	< 430	J	P
Benzyl alcohol	100-51-6	92	U	470	U	12	U	< 570		N
bis(2-Chloroethoxy)methane	111-91-1	1.5	U	6.6	U	6.0	U	< 14		N
bis(2-Chloroethyl)ether	111-44-4	2.0	U	7.3	U	6.4	J	< 16	J	P
bis(2-Ethylhexyl)phthalate	117-81-7	16		100	J	16	J	< 130	J	A
4-Bromophenyl-phenylether	101-55-3	1.4	U	6.6	U	10	J	< 18	J	Р
Butylbenzylphthalate	85-68-7	2.9	U	7.9	U	8.2	J	< 19	J	P
Carbazole	86-74-8	2.0	U	8.4	U	7.2	J	< 18	J	Р
4-Chloro-3-methylphenol	59-50-7	2.6	U	8.1	U	20	U	< 31		N
4-Chloroaniline	106-47-8	3.1	U	79	U	24	U	< 110		N
2-Chloronaphthalene	91-58-7	1.3	U	6.6	U	6.8	J	< 15	J	P
2-Chlorophenol	95-57-8	2.6	U	6.6	U	5.4	J	< 15	J	Р
	7005-72-36		U	6.6	U	9.4	J	< 17	J	P
4-Chlorophenyl phenyl ether	218-01-9	2.3	U	8.4	U	9.9	J	< 21	J	P
Chrysene	210-01-9				-					

## MM-5 Train Summary - Run 2 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-3. HLLWE Run ID: 0010-STRT-2

	CAS Registry	MM-5 Tr Front H Composi (µg)	alf	MM-5 Tr Back Ha Composi (μg)	alf te <sup>2</sup>	MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific Flag <sup>7</sup>
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
								. 110	т	n
Di-n-butylphthalate	84-74-2	1.9	U	100	U	8.4	J	< 110	J	P
Di-n-octylphthalate	117-84-0	5.5	J	150	U	7.6	U	< 160	J	P
Dibenz(a,h)anthracene	53-70-3	5.2	U	160	U	8.4	J	< 170	J	P
Dibenzofuran	132-64-9	1.4	U	6.6	U	8.7	J	< 17	J	P
1,2-Dichlorobenzene	95-50-1	2.2	U	6.8	U	6.6	J	< 16	J	P
1,3-Dichlorobenzene	541-73-1	3.1	U	7.3	U	6.5	J	< 17	J	Р
1,4-Dichlorobenzene	106-46-7	2.9	U	6.8	J	6.2	<u>J</u>	< 16	J	P
3,3'-Dichlorobenzidine	91-94-1	7.1	U	97	U	23	U .	< 130		N
2,4-Dichlorophenol	120-83-2	3.9	U	6.6	U	7.1	U	< 18		N
Diethylphthalate	84-66-2	3.9	U	9.4	U	8.9	J	< 22	J	P
Dimethyl phthalate	131-11-3	1.7	U	6.6	U	8.1	J	< 16	J	P
2,4-Dimethylphenol	105-67-9	7.6	U	50	U	4.7	U	< 62		N
4,6-Dinitro-2-methylphenol	534-52-1	13	U	120	U	4.5	U	< 140		N
2,4-Dinitrophenol	51-28-5	15	U	250	U	12	U	< 280		N
2,4-Dinitrotoluene	121-14-2	4.2	U	6.6	U	8.1	J	< 19	J	P
2,6-Dinitrotoluene	606-20-2	3.4	U	6.6	U	6.3	J	< 16	J	P
1,2-Diphenylhydrazine	122-66-7	1.7	U	6.6	U	7.5	J	< 16	J	P
Fluoranthene	206-44-0	1.3	U	7.1	U	8.5	J	< 17	J	P
Fluorene	86-73-7	1.3	U	6.6	U	8.4	J	< 16	J	P
Hexachlorocyclopentadiene	77-47-4	26	U	130	U	20	U	< 180		P
Hexachlorobenzene	118-74-1	1.5	U	6.6	U	8.3	J	< 16	J	P
Hexachlorobutadiene	87-68-3	3.7	U	9.7	U	6.5	J	< 20	J	P
Hexachloroethane	67-72-1	6.6	U	7.1	U	6.4	J	< 20	J	P
Indeno(1,2,3-cd)pyrene	193-39-5	5.5	U	140	U	7.0	J	< 150	J	P
Isophorone	78-59-1	1.7	U	6.6	U	7.4	J	< 16	J	P
2-Methylnaphthalene	91-57-6	1.5	U	6.6	U	6.8	J	< 15	J	P
2-Methylphenol	95-48-7	6.0	U	39	U	6.3	U	< 51		N
3-Methylphenol & 4-Methylphenol	65794-96-9	6.0	U	26	U	6.8	U	< 39		N

## MM-5 Train Summary - Run 2 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-3. HLLWE Run ID: 0010-STRT-2

	CAS Registry	Front H Compos	MM-5 Train Front Half Composite <sup>1</sup> (μg)		rain alf ite <sup>2</sup>	MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total µg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	(μg) Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
N-Nitroso-di-n-propylamine	621-64-7	1.9	U	6.6	U	6.8	U	< 15		N
N-Nitrosodimethylamine	62-75-9	1.9	U	6.6	U	6.6	J	< 15	J	P
N-Nitrosodiphenylamine	86-30-6	1.6	U	12	U	7.6	J	< 21	J	P
Naphthalene	91-20-3	1.3	U	7.9	U	7.4	J	< 17	J	P
2-Nitroaniline	88-74-4	1.5	U	6.6	U	8.4	U	< 17		N
3-Nitroaniline	99-09-2	10	U	26	U	14	U	< 50	,	N
4-Nitroaniline	100-01-6	6.0	U	26	U	12	U	< 44		N
Nitrobenzene	98-95-1	1.9	U	9.2	J	8.8	J	< 20	J	Р
2-Nitrophenol	88-75-5	8.4	U	21	J	7.9	J	< 37	J	P
4-Nitrophenol	100-02-7	8.7	U	42	U	12	U	< 63		N
2,2'-Oxybis(1-chloropropane) 8	108-60-1	2.6	U	10	U	9.7	J	< 22	J	P
Pentachlorobenzene	608-93-5	1.4	U	6.6	U	6.8	U	< 15		N
Pentachloronitrobenzene	82-68-8	2.0	U	6.6	U	7.9	U	< 16		N
Pentachlorophenol	87-86-5	50	U	250	U	10	U	< 310		N
Phenanthrene	85-01-8	1.3	U	6.6	U	8.6	J	< 17	J	Р
Phenol	108-95-2	2.9	U	23	J	7.4	J	< 33	J	P
Pyrene	129-00-0	1.9	U	6.8	U	9.4	J	< 18	J	P
Pyridine	110-86-1	2.3	U	9.7	U	16	U	< 28		N
1,2,4,5-Tetrachlorobenzene	95-94-3	2.3	U	6.6	U	6.6	U	< 16		N
1,2,4-Trichlorobenzene	120-82-1	1.9	U	7.9	U	7.4	J	< 17	J	P
2,4,5-Trichlorophenol	95-95-4	6.0	U	17	U	6.3	U	< 29		N
2,4,6-Trichlorophenol	88-06-2	3.7	U	10	U	7.6	U_	< 21		N

## MM-5 Train Summary - Run 2 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-3. HLLWE Run ID: 0010-STRT-2

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (µg)	MM-5 Train Back Half Composite <sup>2</sup> (μg)	MM-5 Train Condensate Composite <sup>3</sup> (μg)	MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result Flag	Risk Result Flag <sup>5</sup>	Risk Result Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
TICs <sup>9</sup>							
Furan, 2,5-dimethyl-	625-86-5	9.8			9.8	N,J,M	P
3-Hexanone	589-38-8		96		96	N,J,M	P
Heptane, 2,5-dimethyl-	2216-30-0	7.1			7.1	N,J,M	Р
Benzaldehyde	100-52-7		670		670	N,J,M	P
Formic acid, phenylmethyl este	104-57-4		52		52	N,J,M	P
Dodecane	112-40-3		55		55	N,J,M	P
Tridecane	629-50-5		18		18	N,J,M	Р
Naphthalene, 1-methyl-	90-12-0			7.8	7.8	N,J,M	Р
Tetradecane	629-59-4		56		56	N,J,M	Р
Cyclododecane	294-62-2	5.6			5.6	N,J,M	P
Hexanedioic acid, bis(2-ethylh)	103-23-1			10	10	N,J,M	P
1,2-Benzenedicarboxylic acid,	1330-96-7			8.4	8.4	N,J,M	P
Benzo(e)pyrene	192-97-2			14	14	N,J,M	P

#### Footnotes:

- The MM-5 Train Front Half Composite consists of the Particulate Filter and the Front Half of the Filter Holder and Probe Solvent Rinses.
- The MM-5 Train Back Half Composite consists of the XAD-2 Resin Tube and the Back Half of the Filter Holder and Coil Condenser Solvent Rinses.
- <sup>3</sup> The MM-5 Train Condensate Composite consists of the Condensate and Impinger Contents and the Glassware Solvent Rinses.
- The total mass for each semivolatile compound found in the MM-5 sampling train consists of the sum of the MM-5 train's Front Half Composite contents, the train's Back Half Composite contents, and the Condensate Composite. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) + (Concentration in the Condensate Composite x Condensate Composite Volume) = Total  $\mu g$  in the MM-5 Sampling Train.

Therefore:  $(\mu g) + (\mu g) + (\mu g/Liter \times Liter) = Total \mu g$ 

The MM-5 Train Run Total (in Total  $\mu$ g) is the sum of results for the three (3) MM-5 train sample fractions using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- ♦ When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MM-5 Train Total are the cumulative set of flags for each train component included as part of the MM-5 train total. A flag attached to an MM-5 train component is carried through to the "MM-5 Train Total" column when the associated component analytical result is a significant number in comparison to the MM-5 Train Total. That is, if the MM-5 Train Total is affected by an MM-5 train component analytical result, the flag is carried through to the MM-5 Train Total, but if the MM-5 Train Total is not affected by an MM-5 train component, the flag is not carried through to the MM-5 Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "J" qualifier indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" qualifier indicates that this compound was found in the associated laboratory method blank. Under these conditions these
    values are regarded as estimated values.
  - A "D" qualifier indicates that this result was obtained through dilution of the sample. This original analysis yielded a result that exceeded the calibration range.
  - An "N" qualifier indicates that this compound is a tentatively identified compound (TIC). Therefore the value is estimated.
  - An "E" qualifier indicates that this compound exceeded the calibration range of the instrument.
  - An "A" qualifier indicates that this result is an Aldol-condensation product.
  - An "M" qualifier indicates that this result was measured against the nearest internal standard and assumed a response factor of
    one (1).
  - A "Q" qualifier indicates that this result was quantitated against the response factor of a calibration standard.
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- <sup>8</sup> Bis(2-chloroisopropyl)ether and 2,2'-Oxybis(1-chloropropane) are synonyms.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data. It should be noted that TICs that give the same mass spectral match for GC peaks at different retention times are listed separately with the same compound identity. Under these conditions the compounds are likely indistinguishable isomers of the same compound. However, insufficient evidence is available to determine unequivocal identities.

#### MM-5 Train Summary - Run 4 Train Totals Semivolatile Organic Compounds Analytical Results Summary Table A-4. HLLWE Run ID: 0010-END-2

Field Sample Name:

MM-5 Train

Sample Description:

MM-5 Train Totals for Semivolatile Organic Compounds Analysis

	CAS Registry	MM-5 Ti Front H Compos (µg)	alf	MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total µg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
Target Compound List				-			······································			
Acenaphthene	83-32-9	1.3	U	6.6	U	1.9	U	< 9.8		N
Acenaphthylene	208-96-8	1.3	U	6.6	U	1.6	U	< 9.5		N
Acetophenone	9[8-86-2	2.0	U	32	J	2.6	U	< 37	J	P
Aniline	62-53-3	2.5	U	94	U	18	U	< 110		N
Anthracene	120-12-7	1.3	U	6.6	U	1.6	U	< 9.5	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N
Benzidine	92-87-5	100	U	500	U	66	U	< 670		N
Benzoic acid	65-85-0	100	U	2,600	Е	9.4	U	< 2,700	Е	Р
Benzo(a)anthracene	56-55-3	2.2	U	7.6	U	1.7	U	< 12	/4414444444	N
Benzo(a)pyrene	50-32-8	2.6	U	130	U	1.8	U	< 130		N
Benzo(b)fluoranthene	205-99-2	3.7	U	290	U	4.2	U	< 300		N
Benzo(g,h,i)perylene	191-24-2	7.3	U	160	U	2.1	U	< 170		N
Benzo(k)fluoranthene	207-08-9	5.5	U	420	U	2.9	U	< 430		N
Benzyl alcohol	100-51-6	92	U	470	U	4.2	U	< 570		N
bis(2-Chloroethoxy)methane	111-91-1	1.5	U	6.6	U	2.0	U	< 10		N
bis(2-Chloroethyl)ether	111-44-4	2.0	U	7.3	U	1.7	U	< 11		N
bis(2-Ethylhexyl)phthalate	117-81-7	57		100	J	6.1	J	< 160	J	A
4-Bromophenyl-phenylether	101-55-3	1.4	U	6.6	U	1.4	U	< 9.4		N
Butylbenzylphthalate	85-68-7	2.9	U	7.9	U	2.3	U	< 13		N
Carbazole	86-74-8	2.0	U	8.4	U	2.2	U	< 13		N
4-Chloro-3-methylphenol	59-50-7	2.6	U	8.1	U	6.6	U	< 17		N
4-Chloroaniline	106-47-8	3.1	U	79	U	7.9	U	< 90		N
2-Chloronaphthalene	91-58-7	1.3	U	6.6	U	1.4	U	< 9.3		N
2-Chlorophenol	95-57-8	2.6	U	6.6	U	1.7	U	< 11	<u> </u>	N
4-Chlorophenyl phenyl ether	7005-72-36	1.3	U	6.6	U	3.1	U	< 11		N
Chrysene	218-01-9	2.3	U	8.4	U	1.3	U	< 12		N

## MM-5 Train Summary - Run 4 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-4. HLLWE Run ID: 0010-END-2

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (µg)		MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup>		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
HELIPPIN THE				100		2.2	т т	< 100	J	P
Di-n-butylphthalate	84-74-2	1.9	J	100	U	2.3	U			P
Di-n-octylphthalate	117-84-0	5.5	J	150	U	2.5	J	< 160	J	N N
Dibenz(a,h)anthracene	53-70-3	5.2	U	160	U	2.9	U	< 170		
Dibenzofuran	132-64-9	1.4	U	6.6	U	2.9	U	< 11		N
1,2-Dichlorobenzene	95-50-1	2.2	U	6.8	U	1.7	U	< 11		N
1,3-Dichlorobenzene	541-73-1	3.1	U	7.3	U	1.4	U	< 12		N
1,4-Dichlorobenzene	106-46-7	2.9	U	16	J	2.0	U	< 21	J	P
3,3'-Dichlorobenzidine	91-94-1	7.1	U	97	U	7.9	U	< 110		N
2,4-Dichlorophenol	120-83-2	3.9	U	6.6	U	2.3	U	< 13		N
Diethylphthalate	84-66-2	3.9	U	9.4	U	1.4	U	< 15		N
Dimethyl phthalate	131-11-3	1.7	U	6.6	U	1.3	U	< 9.6		N
2,4-Dimethylphenol	105-67-9	7.6	U	50	U	1.5	U	< 59		N
4,6-Dinitro-2-methylphenol	534-52-1	13	U	120	U	1.5	U	< 130		N
2,4-Dinitrophenol	51-28-5	16	U	250	U	3.9	U	< 270		N
2,4-Dinitrotoluene	121-14-2	4.2	U	6.6	U	2.6	U	< 13		N
2,6-Dinitrotoluene	606-20-2	3.4	U	6.6	U	2.1	U	< 12		N
1,2-Diphenylhydrazine	122-66-7	1.7	U	6.6	U	1.5	U	< 9.8		N
Fluoranthene	206-44-0	1.3	U	7.1	U	1.8	U	< 10		N
Fluorene	86-73-7	1.3	U	6.6	U	2.6	U	< 10		N
Hexachlorocyclopentadiene	77-47-4	26	U	130	U	6.6	U	< 160	1	N
Hexachlorobenzene	118-74-1	1.5	U	6.6	U	2.6	U	< 11		N
Hexachlorobutadiene	87-68-3	3.7	U	9.7	U	1.9	U	< 15		N
Hexachloroethane	67-72-1	6.6	U	7.1	U	1.9	U	< 16		N
Indeno(1,2,3-cd)pyrene	193-39-5	5.5	U	140	U	2.3	U	< 150		N
Isophorone	78-59-1	1.7	U	6.6	U	1.8	U	< 10		N
2-Methylnaphthalene	91 <b>-</b> 57-6	1.5	U	6.6	U	2.3	U	< 10		N
2-Methylphenol	95-48-7	6.0	U	39	U	2.1	U	< 47		N
3-Methylphenol & 4-Methylphenol	65794-96-9	6.0	U	26	U	2.2	U	< 34		N

## MM-5 Train Summary - Run 4 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-4. HLLWE Run ID: 0010-END-2

	CAS Registry	Front H Composi	MM-5 Train Front Half Composite <sup>1</sup> (μg)		rain alf ite <sup>2</sup>	MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	(μg) Risk Result	Flag <sup>5</sup>		Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
N-Nitroso-di-n-propylamine	621-64-7	1.9	U	6.6	U	2.3	U	< 11		N
N-Nitrosodimethylamine	62-75-9	1.9	U	6.6	U	2.2	U	< 11		N
N-Nitrosodiphenylamine	86-30-6	1.6	U	12	U	1.4	U	< 15		N
Naphthalene	91-20-3	1.3	U	7.9	U	1.8	U	< 11		N
2-Nitroaniline	88-74-4	1.5	U	6.6	U	2.9	U	< 11		N
3-Nitroaniline	99-09-2	10	U	26	U	4.7	U	< 41		N
4-Nitroaniline	100-01-6	6.0	U	26	U	3.9	U	< 36		N
Nitrobenzene	98-95-1	1.9	U	8.5	J	1.7	U	< 12	J	Р
2-Nitrophenol	88-75-5	8.4	U	40	J	2.6	U	< 51	J	P
4-Nitrophenol	100-02-7	8.7	U	42	J	3.9	U	< 55	J	Р
2,2'-Oxybis(1-chloropropane) 8	108-60-1	2.6	U	10	U	1.8	U	< 14		N
Pentachlorobenzene	608-93-5	1.4	U	6.6	U	2.3	U	< 10		N
Pentachloronitrobenzene	82-68-8	2.0	U	6.6	U	2.6	U	< 11		N
Pentachlorophenol	87-86-5	50	U	320	U	3.4	U	< 370	ļ	N
Phenanthrene	85-01-8	1.3	U	6.6	U	1.9	U	< 9.8	ļ	N
Phenol	108-95-2	2.9	U	34	J	2.2	J	< 39	J	P
Pyrene	129-00-0	1.9	U	6.8	U	1.4	U	< 10		N
Pyridine	110-86-1	2.3	U	9.7	U	5.2	U	< 17		N
1,2,4,5-Tetrachlorobenzene	95-94-3	2.3	U	6.6	U	2.2	U	< 11		N
1,2,4-Trichlorobenzene	120-82-1	1.9	U	7.9	U	2.2	U	< 12		N
2,4,5-Trichlorophenol	95-95-4	6.0	U	17	U	2.1	U	< 25		N
2,4,6-Trichlorophenol	88-06-2	3.7	U	10	U	2.5	U	< 16		N

### MM-5 Train Summary - Run 4 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-4. HLLWE Run ID: 0010-END-2

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (μg)	MM-5 Train Back Half Composite <sup>2</sup> (μg)	MM-5 Train Condensate Composite <sup>3</sup> (μg)	MM-5 Tot (Tota		Project Specific
Analyte	Number	Risk Result Flag <sup>5</sup>	Risk Result Flag <sup>5</sup>		Total <sup>6</sup>	Flag	Flag <sup>7</sup>
TICs <sup>9</sup>							
3-Hexanone	589-38-8		70		70	N,J,M	P
Benzaldehyde	100-52-7		730		730	N,J,M	Р
2-Cyclohexene-1-one, 3-methyl-	1193-18-6			3.2	3.2	N,J,M	P
Formic acid, phenylmethyl ester	104-57-4		95		95	N,J,M	P
Benzaldehyde, ethyl-	53951-50-1		61		61	N,J,M	Р
Dodecane	112-40-3		37		37	N,J,M	Р
Tridecane	629-50-5		21		21	N,J,M	Р
2,4-Hexadiene	592-46-1		28		28	N,J,M	P
2,5-Diethylphenol	876-20-0		100		100	N,J,M	P
Tetradecane	629-59-4		99		99	N,J,M	P
Hexatriacontane	630-06-8	5.7			5.7	N,J,M	P
Phosphoric acid tributyl ester	126-73-8	24			24	N,J,M	P
Cyclododecane	294-62-2	13			13	N,J,M	Р
Pentadecane	629-62-9	4.5			4.5	N,J,M	P
Heneicosane	629-94-7	9.1			9.1	N,J,M	P
Tetracosane	646-31-1	19			19	N,J,M	P
Pentacosane	629-99-2	35			35	N,J,M	P
Hexacosane	630-01-3	64			64	N,J,M	P
Heptacosane	593-49-7	83			83	N,J,M	P
Pentacosane	629-99-2			2.2	2.2	N,J,M	Р
Hexatriacontane	630-06-8	100			100	N,J,M	Р
Hexatriacontane	630-06-8	67			67	N,J,M	Р
Hexatriacontane	630-06-8	32			32	N,J,M	Р
Eicosane	112-95-8	19			19	N,J,M	P
Tetracosane	646-31-1	7.4			7.4	N,J,M	P

#### Footnotes:

- The MM-5 Train Front Half Composite consists of the Particulate Filter and the Front Half of the Filter Holder and Probe Solvent Rinses.
- The MM-5 Train Back Half Composite consists of the XAD-2 Resin Tube and the Back Half of the Filter Holder and Coil Condenser Solvent Rinses.
- <sup>3</sup> The MM-5 Train Condensate Composite consists of the Condensate and Impinger Contents and the Glassware Solvent Rinses.
- The total mass for each semivolatile compound found in the MM-5 sampling train consists of the sum of the MM-5 train's Front Half Composite contents, the train's Back Half Composite contents, and the Condensate Composite. The calculation is as follows:

(Total µg in the Front Half) + (Total µg in the Back Half) + (Concentration in the Condensate Composite x Condensate Composite Volume)

= Total µg in the MM-5 Sampling Train.

Therefore:  $(\mu g) + (\mu g) + (\mu g/Liter \times Liter) = Total \mu g$ 

The MM-5 Train Run Total (in Total  $\mu g$ ) is the sum of results for the three (3) MM-5 train sample fractions using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- ♦ When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MM-5 Train Total are the cumulative set of flags for each train component included as part of the MM-5 train total. A flag attached to an MM-5 train component is carried through to the "MM-5 Train Total" column when the associated component analytical result is a significant number in comparison to the MM-5 Train Total. That is, if the MM-5 Train Total is affected by an MM-5 train component analytical result, the flag is carried through to the MM-5 Train Total, but if the MM-5 Train Total is not affected by an MM-5 train component, the flag is not carried through to the MM-5 Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "J" qualifier indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" qualifier indicates that this compound was found in the associated laboratory method blank. Under these conditions these values are regarded as estimated values.
  - A "D" qualifier indicates that this result was obtained through dilution of the sample. This original analysis yielded a result that exceeded the calibration range.
  - An "N" qualifier indicates that this compound is a tentatively identified compound (TIC). Therefore the value is estimated.
  - An "E" qualifier indicates that this compound exceeded the calibration range of the instrument.
  - An "A" qualifier indicates that this result is an Aldol-condensation product.
  - An "M" qualifier indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- <sup>8</sup> Bis(2-chloroisopropyl)ether and 2,2'-Oxybis(1-chloropropane) are synonyms.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data. It should be noted that TICs that give the same mass spectral match for GC peaks at different retention times are listed separately with the same compound identity. Under these conditions the compounds are likely indistinguishable isomers of the same compound. However, insufficient evidence is available to determine unequivocal identities.

#### MM-5 Blank Train Summary - Run 2 Train Totals Semivolatile Organic Compounds Analytical Results Summary Table A-5. HLLWE Run ID: 0010-BT-1

Field Sample Name:

MM-5 Blank Train

Sample Description:

MM-5 Blank Train Totals for Semivolatile Organic Compounds Analysis

	CAS Registry	Front H Composi (µg)			MM-5 Train Back Half Composite <sup>2</sup> (µg)		MM-5 Train Condensate Composite <sup>3</sup> (µg)		Train als <sup>4</sup> I μg)	Project Specific Flag <sup>7</sup>
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag	Total <sup>6</sup>	Flag	Flag'
Target Compound List										
Acenaphthene	83-32-9	1.3	U	1.3	U	1.9	U	< 4.5		N
Acenaphthylene	208-96-8	1.3	U	1.3	U	1.6	U	< 4.2		N
Acetophenone	9[8-86-2	2.0	U	6.3	J	2.6	U	< 11	J	Р
Aniline	62-53-3	2.5	U	19	U	18	U	< 40		N
Anthracene	120-12-7	1.3	U	1.3	U	1.6	U	< 4.2		N
Benzidine	92-87-5	100	U	100	U	66	U	< 270		N
Benzoic acid	65-85-0	100	U	100	U	9.4	U	< 210		N
Benzo(a)anthracene	56-55-3	2.2	U	1.5	U	1.7	U	< 5.4		N
Benzo(a)pyrene	50-32-8	2.6	U	1.3	U	1.8	U	< 5.7		N
Benzo(b)fluoranthene	205-99-2	3.7	U	2.9	U	4.2	U	< 11		N
Benzo(g,h,i)perylene	191-24-2	7.3	U	1.6	U	2.1	U	< 11		N
Benzo(k)fluoranthene	207-08-9	5.5	U	4.2	U	2.9	U	< 13		N
Benzyl alcohol	100-51-6	92	U	92	U	4.2	U	< 190		N
bis(2-Chloroethoxy)methane	111-91-1	1.5	U	1.3	U	2.0	U	< 4.8		N
bis(2-Chloroethyl)ether	111-44-4	2.0	U	1.5	U	1.7	U	< 5.2		N
bis(2-Ethylhexyl)phthalate	117-81-7	15		99		15	•	130		A
4-Bromophenyl-phenylether	101-55-3	1.4	U	1.3	U	1.4	U	< 4.1		N
Butylbenzylphthalate	85-68-7	2.9	U	1.6	U	2.3	U	< 6.8		N
Carbazole	86-74-8	2.0	U	1.7	U	2.2	U	< 5.9		N
4-Chloro-3-methylphenol	59-50-7	2.6	U	1.6	U	6.6	U	< 11		N
4-Chloroaniline	106-47-8	3.1	U	16	U	7.9	U	< 27		N
2-Chloronaphthalene	91-58-7	1.3	U	1.3	U	1.4	U	< 4.0		N
2-Chlorophenol	95-57-8	2.6	U	1.3	U	1.7	U	< 5.6		N
4-Chlorophenyl phenyl ether	7005-72-36	1.3	U	1.3	U	3.1	U	< 5.7		N
Chrysene	218-01-9	2.3	U	1.7	U	1.3	U	< 5.3	<u> </u>	N

## MM-5 Blank Train Summary - Run 2 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-5. HLLWE Run ID: 0010-BT-1

	CAS Registry	MM-5 Ti Front H Composi (μg)	alf ite <sup>1</sup>	MM-5 Train Back Half Composite <sup>2</sup> (μg)		MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total µg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Risk Result	Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
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Di-n-butylphthalate	84-74-2	1.9	J	20	U	2.3	J	< 24	J	P
Di-n-octylphthalate	117-84-0	5.5	J	10		3.3	J	< 19	J	A
Dibenz(a,h)anthracene	53-70-3	5.2	U	1.6	U	2.9	U	< 9.7		N
Dibenzofuran	132-64-9	1.4	U	1.3	U	2.9	U	< 5.6		N
1,2-Dichlorobenzene	95-50-1	2.2	U	1.3	U	1.7	U	< 5.2		N
1,3-Dichlorobenzene	541-73-1	3.1	U	1.5	U	1.4	U	< 6.0		N
1,4-Dichlorobenzene	106-46-7	2.9	U	12		2.0	U	< 17		Р
3,3'-Dichlorobenzidine	91-94-1	7.1	U	19	U	7.9	U	< 34		N
2,4-Dichlorophenol	120-83-2	3.9	U	1.3	U	2.3	U	< 7.5		N
Diethylphthalate	84-66-2	3.9	U	1.9	U	1.4	U	< 7.2		N
Dimethyl phthalate	131-11-3	1.7	U	1.3	U	1.3	U	< 4.3		N
2,4-Dimethylphenol	105-67-9	7.6	U	10	U	1.5	U	< 19		N
4,6-Dinitro-2-methylphenol	534-52-1	13	U	23	U	1.5	U	< 38		N
2,4-Dinitrophenol	51-28-5	15	U	50	U	3.9	U	< 69		N
2,4-Dinitrotoluene	121-14-2	4.2	U	1.3	U	2.6	U	< 8.1		N
2,6-Dinitrotoluene	606-20-2	3.4	U	1.3	U	2.1	U	< 6.8		Ν
1,2-Diphenylhydrazine	122-66-7	1.7	U	1.3	U	1.5	U	< 4.5		N
Fluoranthene	206-44-0	1.3	U	1.4	U	1.8	U	< 4.5		N
Fluorene	86-73-7	1.3	U	1.3	U	2.6	U	< 5.2		N
Hexachlorocyclopentadiene	77-47-4	26	U	26	U	6.6	U	< 59		N
Hexachlorobenzene	118-74-1	1.5	U	1.3	U	2.6	U	< 5.4		N
Hexachlorobutadiene	87-68-3	3.7	U	1.9	U	1.9	U	< 7.5		N
Hexachloroethane	67-72-1	6.6	U	1.4	U	1.9	U	< 9.9		N
Indeno(1,2,3-cd)pyrene	193-39-5	5.5	U	1.4	U	2.3	U	< 9.2		N
Isophorone	78-59-1	1.7	U	1.3	U	1.8	U	< 4.8		N
2-Methylnaphthalene	91-57-6	1.5	U	1.3	U	2.3	U	< 5.1		N
2-Methylphenol	95-48-7	6.0	U	7.9	U	2.1	U	< 16		N
3-Methylphenol & 4-Methylphenol	65794-96-9	6.0	U	5.2	U	2.2	U	< 13		N

### MM-5 Blank Train Summary - Run 2 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-5. HLLWE Run ID: 0010-BT-1

	CAS Registry	Front H Compos	Front Half Composite <sup>1</sup> (µg)		rain alf ite <sup>2</sup>	MM-5 Train Condensate Composite <sup>3</sup> (μg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific
Analyte	Number	Risk Result	Flag <sup>5</sup>	(μg) Risk Result	Flag <sup>5</sup>		Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
N-Nitroso-di-n-propylamine	621-64-7	1.9	U	1.3	U	2.3	U	< 5.5		N
N-Nitrosodimethylamine	62-75-9	1.9	U	1.3	U	2.2	U	< 5.4		N
N-Nitrosodiphenylamine	86-30-6	1.6	U	2.3	U	1.4	U	< 5.3		N
Naphthalene	91-20-3	1.3	U	1.6	J	1.8	U	< 4.7	J	Р
2-Nitroaniline	88-74-4	1.5	U	1.3	U	2.9	U	< 5.7		N
3-Nitroaniline	99-09-2	10	U	5.2	U	4.7	U	< 20		N
4-Nitroaniline	100-01-6	6.0	U	5.2	U	3.9	U	< 15		N
Nitrobenzene	98-95-1	1.9	U	1.5	U	1.7	U	< 5.1		N
2-Nitrophenol	88-75-5	8.4	U	1.3	U	2.6	U	< 12		N
4-Nitrophenol	100-02-7	8.7	U	8.7	U	3.9	U	< 21		N
2,2'-Oxybis(1-chloropropane) 8	108-60-1	2.6	U	2.0	U	1.8	U	< 6.4		N
Pentachlorobenzene	608-93-5	1.4	U	1.3	U	2.3	U	< 5.0	·	N
Pentachloronitrobenzene	82-68-8	2.0	U	1.3	U	2.6	U	< 5.9		N
Pentachlorophenol	87-86-5	50	U	50	U	3.4	U	< 100		N
Phenanthrene	85-01-8	1.3	U	1.3	U	1.9	U	< 4.5		N
Phenol	108-95-2	2.9	U	2.4	U	2.2	U	< 7.5	·	N
Pyrene	129-00-0	1.9	U	1.4	U	1.4	U	< 4.7		N
Pyridine	110-86-1	2.3	U	1.9	U	5.2	U	< 9.4		N
1,2,4,5-Tetrachlorobenzene	95-94-3	2.3	U	1.3	U	2.2	U	< 5.8		N
1,2,4-Trichlorobenzene	120-82-1	1.9	U	1.5	U	2.2	U	< 5.6		N
2,4,5-Trichlorophenol	95-95-4	6.0	U	3.4	U	2.1	U	< 12	-	N
2,4,6-Trichlorophenol	88-06-2	3.7	U	2.0	U	2.5	U	< 8.2		N

### MM-5 Blank Train Summary - Run 2 Train Totals (Continued) Semivolatile Organic Compounds Analytical Results Summary Table A-5. HLLWE Run ID: 0010-BT-1

	CAS Registry	MM-5 Train Front Half Composite <sup>1</sup> (µg)	Front Half Composite Composite (µg)  Front Half Composite (µg)		MM-5 Train Totals <sup>4</sup> (Total μg)		Project Specific Flag <sup>7</sup>
Analyte	Number	Risk Result Flag <sup>5</sup>	Risk Result Flag <sup>5</sup>	Risk Result Flag <sup>5</sup>	Total <sup>6</sup>	Flag	Flag <sup>7</sup>
TICs9							
Furan, 2,5-dimethyl-	625-86-5	12			12	N,J,M	P
Heptane, 2,5-dimethyl-	2216-30-0	9.0			9.0	N,J,M	P
Heptane, 2,3-dimethyl-	3074-71-3	11			11	N,J,M	Р
Benzaldehyde	100-52-7		7.3		7.3	N,J,M	P
Benzoic acid, methyl ester	93-58-3		8.4		8.4	N,J,M	P
Benzaldehyde, ethyl-	53951-50-1		14		14	N,J,M	P
Pentadecane	629-62-9	4.4			4.4	N,J,M	Р
Cyclododecane	294-62-2	11	8.7	10	30	N,J,M	P
Heptadecane	629-78-7	3.0			3.0	N,J,M	P
Eicosane	112-95-8			3.7	3.7	N,J,M	P
Heneicosane	629-94-7		4.5		4.5	N,J,M	P
Octodecane	593-45-3		10		10	N,J,M	Р
Phosphine oxide, triphenyl-	791-28-6			26	26	N,J,M	P
Nonacosane	630-03-5		10		10	N,J,M	Р
Eicosane	112-95-8		14		14	N,J,M	P
Hexatriacontane	630-06-8		23		23	N,J,M	P
Tetracosane	646-31-1		18		18	N,J,M	Р
Heneicosane	629-94-7			3.6	3.6	N,J,M	Р
Tetratriacontane	14167-59-0		11		11	N,J,M	Р
Eicosane	112-95-8		9.2		9.2	N,J,M	Р

#### Footnotes:

- The MM-5 Train Front Half Composite consists of the Particulate Filter and the Front Half of the Filter Holder and Probe Solvent Rinses.
- The MM-5 Train Back Half Composite consists of the XAD-2 Resin Tube and the Back Half of the Filter Holder and Coil Condenser Solvent Rinses.
- <sup>3</sup> The MM-5 Train Condensate Composite consists of the Condensate and Impinger Contents and the Glassware Solvent Rinses.
- The total mass for each semivolatile compound found in the MM-5 sampling train consists of the sum of the MM-5 train's Front Half Composite contents, the train's Back Half Composite contents, and the Condensate Composite. The calculation is as follows:

(Total µg in the Front Half) + (Total µg in the Back Half) + (Concentration in the Condensate Composite x Condensate Composite Volume)

= Total  $\mu g$  in the MM-5 Sampling Train.

Therefore:  $(\mu g) + (\mu g) + (\mu g/Liter \times Liter) = Total \mu g$ 

The MM-5 Train Run Total (in Total  $\mu g$ ) is the sum of results for the three (3) MM-5 train sample fractions using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MM-5 Train Total are the cumulative set of flags for each train component included as part of the MM-5 train total. A flag attached to an MM-5 train component is carried through to the "MM-5 Train Total" column when the associated component analytical result is a significant number in comparison to the MM-5 Train Total. That is, if the MM-5 Train Total is affected by an MM-5 train component analytical result, the flag is carried through to the MM-5 Train Total, but if the MM-5 Train Total is not affected by an MM-5 train component, the flag is not carried through to the MM-5 Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- 5 This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "J" qualifier indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" qualifier indicates that this compound was found in the associated laboratory method blank. Under these conditions these values are regarded as estimated values.
  - A "D" qualifier indicates that this result was obtained through dilution of the sample. This original analysis yielded a result that exceeded the calibration range.
  - An "N" qualifier indicates that this compound is a tentatively identified compound (TIC). Therefore the value is estimated.
  - An "E" qualifier indicates that this compound exceeded the calibration range of the instrument.
  - ♦ An "A" qualifier indicates that this result is an Aldol-condensation product.
  - An "M" qualifier indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- <sup>8</sup> Bis(2-chloroisopropyl)ether and 2,2'-Oxybis(1-chloropropane) are synonyms.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data. It should be noted that TICs that give the same mass spectral match for GC peaks at different retention times are listed separately with the same compound identity. Under these conditions the compounds are likely indistinguishable isomers of the same compound. However, insufficient evidence is available to determine unequivocal identities.

### MM-5 Train Analytical Results Summary Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank

Field Sample Name:

MM-5 Train XAD-2 Resin Tube Trip Blank/Reagent Blank

Sample Description:

MM-5 Train XAD-2 Resin Tube Trip Blank/Reagent Blank for Semivolatile Organic Compounds Analysis

Field Sample ID:

A-3378

STL Sample No.

H1F250162-007

	CAS Registry		XAD-2 R	MM-5 esin Tube Tri (με	p Blank/Re	agent Blank	
Analyte	Number	Lab Result <sup>2</sup>	MDL <sup>3</sup>	RDL⁴	RL <sup>5</sup>	Risk Result	Flag <sup>6</sup>
Acenaphthene	83-32-9	ND	0.50	1.3	10	< 1.3	
Acenaphthylene	208-96-8	ND	0.50	1.3	10	< 1.3	
Acetophenone	98-86-2	2.7	2.4	6.3	10	6.3	J
Aniline	62-53-3	ND	7.3	19	20	< 19	
Anthracene	120-12-7	ND	0.50	1.3	10	< 1.3	
Benzidine	92-87-5	ND	51	130	100	< 100	
Benzoic acid	65-85-0	ND	46	120	100	< 100	
Benzo(a)anthracene	56-55-3	ND	0.58	1.5	10	< 1.5	
Benzo(a)pyrene	50-32-8	ND	0.50	1.3	10	< 1.3	
Benzo(b)fluoranthene	205-99-2	ND	1.1	2.9	10	< 2.9	
Benzo(g,h,i)perylene	191-24-2	ND	0.62	1.6	10	< 1.6	
Benzo(k)fluoranthene	207-08-9	ND	1.6	4.2	10	< 4.2	
Benzyl alcohol	100-51-6	ND	35	92	100	< 92	
bis(2-Chloroethoxy)methane	111-91-1	ND	0.50	1.3	10	< 1.3	
bis(2-Chloroethyl)ether	111-44-4	ND	0.56	1.5	10	< 1.5	**************************************
bis(2-Ethylhexyl)phthalate	117-81-7	ND	10	26	20	< 20	
4-Bromophenyl-phenylether	101-55-3	ND	0.50	1.3	10	< 1.3	
Butylbenzylphthalate	85-68-7	ND	0.61	1.6	10	< 1.6	
Carbazole	86-74-8	ND	0.64	1.7	10	< 1.7	
4-Chloro-3-methylphenol	59-50-7	ND	0.62	1.6	10	< 1.6	
4-Chloroaniline	106-47-8	ND	6.0	16	20	< 16	
2-Chloronaphthalene	91-58-7	ND	0.50	1.3	10	< 1.3	***************************************
2-Chlorophenol	95-57-8	ND	0.50	1.3	10	< 1.3	
4-Chlorophenyl phenyl ether	7005-72-36	ND	0.50	1.3	10	< 1.3	
Chrysene	218-01-9	ND	0.64	1.7	10	< 1.7	

### MM-5 Train Analytical Results Summary (Continued) Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank

	CAS Registry	stry (µg) <sup>1</sup>											
Analyte	Number	Lab Result <sup>2</sup>	MDL <sup>3</sup>	RDL <sup>4</sup>	RL <sup>5</sup>	Risk Result	Flag <sup>6</sup>						
Di-n-butylphthalate	84-74-2	ND	10	26	20	< 20							
Di-n-octylphthalate	117-84-0	ND	0.56	1.5	10	< 1.5	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,						
Dibenz(a,h)anthracene	53-70-3	ND	0.60	1.6	10	< 1.6							
Dibenzofuran	132-64-9	ND	0.50	1.3	10	< 1.3							
1,2-Dichlorobenzene	95-50-1	ND	0.51	1.3	10	< 1.3							
1,3-Dichlorobenzene	541-73-1	ND	0.57	1.5	10	< 1.5							
1,4-Dichlorobenzene	106-46-7	9.0	0.53	1.4	10	9.0	J						
3,3'-Dichlorobenzidine	91-94-1	ND	7.4	19	50	< 19							
2,4-Dichlorophenol	120-83-2	ND	0.50	1.3	10	< 1.3							
Diethylphthalate	84-66-2	ND	0.73	1.9	10	< 1.9							
Dimethyl phthalate	131-11-3	ND	0.50	1.3	10	< 1.3							
2,4-Dimethylphenol	105-67-9	ND	6.3	16	10	< 10							
4,6-Dinitro-2-methylphenol	534-52-1	ND	8.7	23	50	< 23							
2,4-Dinitrophenol	51-28-5	ND	22	58	50	< 50							
2,4-Dinitrotoluene	121-14-2	ND	0.50	1.3	10	< 1.3	,,						
2,6-Dinitrotoluene	606-20-2	ND	0.50	1.3	10	< 1.3							
1,2-Diphenylhydrazine	122-66-7	ND	0.50	1.3	10	< 1.3							
Fluoranthene	206-44-0	ND	0.54	1.4	10	< 1.4							
Fluorene	86-73-7	ND	0.50	1.3	10	< 1.3							
Hexachlorocyclopentadiene	77-47-4	ND	10	26	50	< 26							
Hexachlorobenzene	118-74-1	ND	0.50	1.3	10	< 1.3							
Hexachlorobutadiene	87-68-3	ND	0.74	1.9	10	< 1.9							
Hexachloroethane	67-72-1	ND	0.54	1.4	10	< 1.4							
Indeno(1,2,3-cd)pyrene	193-39-5	ND	0.54	1.4	10	< 1.4							
Isophorone	78-59-1	ND	0.50	1.3	10	< 1.3							
2-Methylnaphthalene	91-57-6	ND	0.50	1.3	10	< 1.3							
2-Methylphenol	95-48-7	ND	3.0	7.9	10	< 7.9							
3-Methylphenol & 4-Methylphenol	65794-96-9	ND	2.0	5.2	10	< 5.2							

### MM-5 Train Analytical Results Summary (Continued) Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank

	CAS Registry	MM-5 Train XAD-2 Resin Tube Trip Blank/Reagent Blank (μg) <sup>1</sup>											
Analyte	Number	Lab Result <sup>2</sup>	MDL <sup>3</sup>	RDL <sup>4</sup>	RL <sup>5</sup>	Risk Result	Flag <sup>6</sup>						
N-Nitroso-di-n-propylamine	621-64-7	ND	0.50	1.3	10	< 1.3							
N-Nitrosodimethylamine	62-75-9	ND	0.50	1.3	10	< 1.3							
N-Nitrosodiphenylamine	86-30-6	ND	0.87	2.3	10	< 2.3							
Naphthalene	91-20-3	ND	0.60	1.6	10	< 1.6							
2-Nitroaniline	88-74-4	ND	0.50	1.3	50	< 1.3							
3-Nitroaniline	99-09-2	ND	2.0	5.2	50	< 5.2							
4-Nitroaniline	100-01-6	ND	2.0	5.2	50	< 5.2							
Nitrobenzene	98-95-1	ND	0.57	1.5	10	< 1.5							
2-Nitrophenol	88-75-5	ND	0.50	1.3	10	< 1.3							
4-Nitrophenol	100-02-7	ND	3.3	8.7	50	< 8.7							
2,2'-Oxybis(1-chloropropane) 7	108-60-1	ND	0.76	2.0	10	< 2.0							
Pentachlorobenzene	608-93-5	ND	0.50	1.3	10	< 1.3							
Pentachloronitrobenzene	82-68-8	ND	0.50	1.3	50	< 1.3							
Pentachlorophenol	87-86-5	ND	25	66	50	< 50							
Phenanthrene	85-01-8	ND	0.50	1.3	10	< 1.3							
Phenol	108-95-2	ND	0.90	2.4	10	< 2.4							
Pyrene	129-00-0	ND	0.53	1.4	10	< 1.4							
Pyridine	110-86-1	ND	0.74	1.9	20	< 1.9							
1,2,4,5-Tetrachlorobenzene	95-94-3	ND	0.50	1.3	10	< 1.3							
1,2,4-Trichlorobenzene	120-82-1	ND	0.59	1.5	10	< 1.5							
2,4,5-Trichlorophenol	95-95-4	ND	1.3	3.4	10	< 3.4							
2,4,6-Trichlorophenol	88-06-2	ND	0.75	2.0	10	< 2.0							

### BECHTEL BWXT IDAHO, LLC (BBWI)

INTEC HLLWE Effluent Gas Emissions Inventory

Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### MM-5 Train Analytical Results Summary (Continued) Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank

#### Sampling Surrogate Recoveries:

Sampling Surrogate Compound <sup>8</sup>	Percent Recovery (%)	Project Target Recovery Limits (%)
<sup>13</sup> C <sub>6</sub> -Naphthalene	66%	50-150%

#### Surrogate Standard Recoveries:

Surrogate Standard Compound	Percent Recovery (%)	Laboratory Recovery Limits (%)
	55%	19-100%
2-Fluorophenol	3370	
Phenol- d₅	61%	15-124%
Nitrobenzene-d₅	63%	35-122%
2-Fluorobiphenyl	67%	34-115%
2,4,6-Tribromophenol	0.0%9	33-130%
Terphenyl-d <sub>14</sub>	84%	28-132%

#### Sample Collection and Analysis Dates:

Date(s) Collected:

June 21, 2001

Date(s) of Extraction:

June 26, 2001

Date(s) of Analysis:

July 02, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0010:

"Modified Method 5 Sampling Train"

SW-846 Method 3542:

"Extraction of Semivolatile Analytes Collected Using Method 0010 (Modified Method 5 Sampling Train)"

SW-846 Method 8270C:

"Semivolatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)

Capillary Column Technique"

### MM-5 Train Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank Tentatively Identified Compound (TIC) Summary

TICs <sup>10</sup>	CAS Registry Number	Approximate Retention Time (min.)	Sample Result (µg)	TIC Flag
Benzaldehyde	100-52-7	3.60	5.3	N,J,M
Benzoic acid, methyl ester	93-58-3	4.72	6.5	N,J,M
Benzaldehyde, ethyl-	53951-50-1	5.40	4.6	N,J,M
Heptacosane	593-49-7	11.48	3.1	N,J,M
Heneicosane	629-94-7	11.73	5.8	N,J,M
Tetratracontane	7098-22-8	11.96	8.5	N,J,M
Hexatriacontane	630-06-8	12.19	8.9	N,J,M
Hexatriacontane	630-06-8	12.41	8.2	N,J,M
Heneicosane	629-94-7	12.64	14	N,J,M
Heptacosane	593-49-7	12.90	9.1	N,J,M

### MM-5 Train Analytical Results Summary Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank (Continued)

#### Footnotes:

- Based on the selection rules, the **bolded** value is the value or default value assigned to the analyte using the following guidelines:
  - When the analytical result is greater than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the result selected by boldface type is the RDL.
  - When the analytical result is not detected down to the MDL, the result selected by boldface type is the RDL.
  - ♦ It should be noted that when the RDL is selected using the guidelines above, but the RL is less than the RDL, the RL is included as the "Risk Result".
- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- <sup>5</sup> The RL is the laboratory Reporting Limit (RL).
- <sup>6</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "J" qualifier indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" qualifier indicates that this compound was found in the associated laboratory method blank. Under these conditions these values are regarded as estimated values.
  - A "D" qualifier indicates that this result was obtained through dilution of the sample. This original analysis yielded a result that exceeded the calibration range.
  - An "N" qualifier indicates that this compound is a tentatively identified compound (TIC). Therefore the value is estimated.
  - An "E" qualifier indicates that this compound exceeded the calibration range of the instrument.

### MM-5 Train Analytical Results Summary Table A-6. Run 2, XAD-2 Resin Tube Trip Blank/Reagent Blank (Continued)

- An "A" qualifier indicates that this result is an Aldol-condensation product.
- An "M" qualifier indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- Bis(2-chloroisopropyl)ether and 2,2'-Oxybis(1-chloropropane) are synonyms.
- This material is a sampling surrogate and is an isotopically-labeled compound spiked on the XAD-2 Resin Tube prior to the collection of sample on the MM-5 sampling train.
- <sup>9</sup> This percent recovery is outside of the laboratory target recovery range.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data. It should be noted that TICs that give the same mass spectral match for GC peaks at different retention times are listed separately with the same compound identity. Under these conditions the compounds are likely indistinguishable isomers of the same compound. However, insufficient evidence is available to determine unequivocal identities.

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### BECHTEL BWXT IDAHO, LLC (BBWI) INTEC HLLWE Effluent Gas Emissions Inventory Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### **VOST Summary - Run 1 Train Totals** Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-7. HLLWE Run ID: 0031-STRT-1

Field Sample Name: Sample Description: Volatile Organic Sampling Train (VOST) Totals
Sample Description: Tenax® and Anasorb 747® Tube Sets (Sets #1, #2, #3, and #4) and the VOST Condensate for Volatile Organic Compounds (VOC) Analysis

		VOS Tube So (Total pa	et #1	VOS Tube Se (Total µ	et #2	VOS Tube Se (Total με	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Tota	nsate	VOS Tota (Total	l¹	
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
Target Compound List					.,,,,,,									
Acetone	67 <b>-</b> 64-1	2.4	В	1.1	В	1.0	В	1.3	В	1.4	В	7.2	В	A
Acrylonitrile	107-13-1	< 0.58		< 0.58		< 0.58		< 0.58		< 0.51		< 2.8	***************	N
Benzene	71-43-2	< 0.17		< 0.11		< 0.046		< 0.043		< 0.027		< 0.40		Р
Bromobenzene	108-86-1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.036		< 0.12		N
Bromochloromethane	74-97-5	< 0.030		< 0.030		< 0.030		< 0.030		< 0.028		< 0.15		N
Bromodichloromethane	75-27-4	< 0.022		< 0.022		< 0.022		< 0.022		< 0.033		< 0.12		N
Bromoform	75-25-2	< 0.038		< 0.038		< 0.038		< 0.038		< 0.025		< 0.18		N
Bromomethane	74-83-9	< 0.030	J	< 0.038	J	< 0.030	J	< 0.054	J	< 0.020	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.17	J	P
2-Butanone	78-93-3	< 0.20	J	< 0.20		< 0.20		< 0.20		< 0.093		< 0.89	J	Р
n-Butylbenzene	104-51-8	< 0.032		< 0.032		< 0.032		< 0.032		< 0.025		< 0.15		N
sec-Butylbenzene	135-98-8	< 0.017		< 0.017		< 0.017		< 0.017		< 0.017		< 0.085		N
tert-Butylbenzene	98-06-6	< 0.032		< 0.032		< 0.032		< 0.032		< 0.014		< 0.14		N
Carbon disulfide	75-15-0	< 0.37		< 0.24		< 0.16		< 0.19		< 0.011		< 0.97		Р
Carbon tetrachloride	56-23-5	< 0.036		< 0.036		< 0.036		< 0.036		< 0.020		< 0.16		N
Chlorobenzene	108-90-7	< 0.017	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.017		< 0.017		< 0.017		< 0.027		< 0.095		N
Chlorodibromomethane	124-48-1	< 0.030		< 0.030		< 0.030		< 0.030		< 0.029		< 0.15		N
Chloroethane	75-00-3	< 0.036		< 0.036	J	< 0.036	J	< 0.043	J	< 0.014		< 0.16	J	P
Chloroform	67-66-3	< 0.036	,	< 0.036		< 0.036	J	< 0.10		< 0.031		< 0.24	J	P
Chloromethane	74-87-3	0.15	J	0.28	. J	0.14	J	0.61	•,,,•,,	< 0.011		1.2	J	P
2-Chlorotoluene	95-49-8	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
4-Chlorotoluene	106-43-4	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
1,2-Dibromo-3-chloropropane	96-12-8	< 0.058		< 0.058		< 0.058		< 0.058		< 0.042		< 0.27		N
1,2-Dibromoethane	106-93-4	< 0.040		< 0.040		< 0.040		< 0.040		< 0.042		< 0.20		N
Dibromomethane	74-95-3	< 0.034		< 0.034		< 0.034		< 0.034		< 0.030		< 0.17		N
1,2-Dichlorobenzene	95-50-1	< 0.040		< 0.040		< 0.040	······································	< 0.040		< 0.019		< 0.18		N
1,3-Dichlorobenzene	541-73-1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.022		< 0.10		N
1,4-Dichlorobenzene	106-46-7	< 0.028		< 0.028		< 0.028		< 0.028		< 0.023		< 0.14		N
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## VOST Summary - Run 1 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-7. HLLWE Run ID: 0031-STRT-1

		VOST Tube Set #1 (Total μg/Set)		VOS Tube S (Total μ	et #2	VOS Tube Se (Total με	et #3	VOS Tube S (Total µ	et #4	VOS Conde (Total	nsate	VOST Total <sup>1</sup> (Total μg)		
Analyte	CAS Registry Number	Risk Result <sup>2</sup>		Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
Dichlorodifluoromethane	75-71-8	< 0.064		< 0.061		< 0.057		< 0.062		< 0.011		< 0.26		P
1,1-Dichloroethane	75-34-3	< 0.034		< 0.034	***	< 0.034		< 0.034		< 0.017		< 0.15		N
1,2-Dichloroethane	107-06-2	< 0.034	J	< 0.034		< 0.034		< 0.034	***************************************	< 0.020		< 0.16	J	Р
1,1-Dichloroethene	75-35-4	< 0.036		< 0.036	J	< 0.036	J	< 0.036	J	< 0.016		< 0.16	J	Р
cis-1,2-Dichloroethene	156-59-2	< 0.032		< 0.032		< 0.032	***************************************	< 0.032		< 0.021		< 0.15		N
trans-1,2-Dichloroethene	156-60-5	< 0.038		< 0.038	······	< 0.038		< 0.038		< 0.014		< 0.17		N
1,2-Dichloropropane	78-87-5	< 0.026		< 0.026		< 0.026		< 0.026		< 0.023		< 0.13		N
1,3-Dichloropropane	142-28-9	< 0.038		< 0.038		< 0.038	***************************************	< 0.038	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.022		< 0.17		N
2.2-Dichloropropane	594-20-7	< 0.036		< 0.036		< 0.036		< 0.036		< 0.011		< 0.16		N
1,1-Dichloropropene	563-58-6	< 0.040		< 0.040		< 0.040		< 0.040		< 0.016		< 0.18		N
cis-1,3-Dichloropropene	10061-01-5	< 0.024		< 0.024		< 0.024		< 0.024		< 0.030	***************************************	< 0.13		N
trans-1,3-Dichloropropene	10061-02-6	< 0.030		< 0.030		< 0.030		< 0.030		< 0.028		< 0.15		N
Ethylbenzene	100-41-4	< 0.018		< 0.018		< 0.018		< 0.018		< 0.021		< 0.093		N
Hexachlorobutadiene	87-68-3	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.22		N
2-Hexanone	591-78-6	< 0.13		< 0.13		< 0.13		< 0.13	•••••	< 0.036		< 0.56		N
Isopropylbenzene	98-82-8	< 0.013		< 0.013		< 0.013		< 0.013		< 0.018		< 0.070		N
p-Isopropyltoluene	99-87-6	< 0.024		< 0.024		< 0.024		< 0.024		< 0.018		< 0.11	•	N
Methylene chloride	75-09-2	13	E	3.0	E,B	1.1	В	0.96	В	0.12	В	18	E,B	A
4-Methyl-2-pentanone	108-10-1	< 0.14		< 0.14		< 0.14		< 0.14		< 0.030		< 0.59		N
Naphthalene	91-20-3	< 0.050		< 0.050		< 0.050		< 0.050		< 0.011		< 0.21		N
n-Propylbenzene	103-65-1	< 0.011		< 0.011		< 0.011		< 0.012		< 0.022		< 0.067		N
Styrene	100-42-5	< 0.014		< 0.014		< 0.014		< 0.014		< 0.022		< 0.078		N
1,1,1,2-Tetrachloroethane	630-20-6	< 0.019		< 0.019		< 0.019		< 0.019		< 0.023		< 0.099		N
1,1,2,2-Tetrachloroethane	79-34-5	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.22		N
Tetrachloroethene	127-18-4	< 0.032		< 0.032		< 0.032		< 0.032		< 0.021		< 0.15		N
Toluene	108-88-3	< 0.032	J	0.041	J	< 0.023	J	0.064	J	< 0.028		< 0.19	J	Р
1.2,3-Trichlorobenzene	87-61-6	< 0.050		< 0.050		< 0.050		< 0.050		< 0.011		< 0.21		N
1,2,4-Trichlorobenzene	120-82-1	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.22		N
1,1,1-Trichloroethane	71-55-6	< 0.044		< 0.044		< 0.044		< 0.044		< 0.018	,,	< 0.19		N
1,1,2-Trichloroethane	79-00-5	< 0.036		< 0.036		< 0.036		< 0.036		< 0.022		< 0.17		N

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### VOST Summary - Run 1 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-7. HLLWE Run ID: 0031-STRT-1

		VOS Tube S (Total μ	et #1	VOS Tube So (Total μ	et #2	VOS Tube Se (Total μ	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Tota	nsate	VOS Tota (Total	al <sup>1</sup>	
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
											***************************************			
Trichloroethene	79-01-6	< 0.034		< 0.034		< 0.034		< 0.034		< 0.020		< 0.16		N
Trichlorofluoromethane	75-69-4	< 0.036	J	< 0.036	J	< 0.036	J	< 0.036	J	< 0.011		< 0.16	J	P
1,2,3-Trichloropropane	96-18-4	< 0.050		< 0.050		< 0.050		< 0.050		< 0.036		< 0.24		N
1,2,4-Trimethylbenzene	95-63-6	< 0.015		< 0.015		< 0.015		< 0.015		< 0.042		< 0.10		N
1,3,5-Trimethylbenzene	108-67-8	< 0.010		< 0.010		< 0.010		< 0.010		< 0.019		< 0.059		N
Vinyl chloride	75-01-4	< 0.013		< 0.016	J	< 0.013	J	< 0.022	J	< 0.068		< 0.13	J	P
m-Xylene & p-Xylene	136777-61-2	< 0.10		< 0.10		< 0.10		< 0.10		< 0.042		< 0.44		N
o-Xylene	95-47-6	< 0.013		< 0.013		< 0.013		< 0.013		< 0.025		< 0.077		N
TICs <sup>7</sup>							***************************************							
Hexane, 2-methyl-	591-76-4		***************************************					0.17				0.17	N,J,M	Р
Pentane, 2,3-dimethyl-	565-59-3							0.18				0.18	N,J,M	Р
Butane, 1-chloro-	109-69-3							0.057				0.057	N,J,M	P
Hexane, 3-methyl-	589-34-4							0.38				0.38	N,J,M	P
Cyclohexene	110-83-8			0.044		0.027		0.033				0.10	N,J,M	P
1-Heptene	592-76-7							0.054				0.054	N,J,M	Р
Cyclohexane, methyl-	108-87-2							0.11				0.11	N,J,M	Р
Hexane, 2,4-dimethyl-	589-43-5							0.11				0.11	N,J,M	Р
Cyclopentane, ethyl-	1640-89-7							0.028				0.028	N,J,M	P
Octane	111-65-9	0.027										0.027	N,J,M	Р
Decane	124-18-5			0.055		0.060			47744444444			0.12	N,J,M	P
Undecane	1120-21-4	0.44		0.23				0.37			.,,	1.0	N,J,M	P
Undecane, 5-methyl-	1632-70-8	0.13		0.091		0.14		0.28				0.64	N,J,M	P
Decane, 2,9-dimethyl-	1002-17-1	0.064										0.064	N,J,M	P
Dodecane	112-40-3	5.3		4.6		7.2		14			.,,,	31	N,J,M	Р
Undecane, 2,6-dimethyl-	17301-23-4							0.11				0.11	N,J,M	P
Cyclohexane, hexyl-	4292-75 <b>-</b> 5				,,,,,	0.059						0.059	N,J,M	Р
Tridecane	629-50-5	0.39		0.32		0.51		1.6				2.8	N,J,M	P
Tetradecane	629-59-4	0.15		0.19		0.22		0.36				0.92	N,J,M	P

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#### Footnotes:

- The Method 0031 VOST Run Total (in Total μg) is the sum of results for the four (4) VOST tube sets and the condensate sample collected during the same sampling run using the following guidelines:
  - When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
  - When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
  - It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the VOST Total are the cumulative set of flags contributed by each train tube set included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Total" column when the associated component analytical result is a significant number in comparison to the VOST Total. That is, if the VOST Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Total, but if the VOST Total is not affected by a VOST component, the associated flag is not carried through to the VOST Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

The Method 0031 VOST Tube Set (Total μg/Set) result consists of the sum of the analytical results for the two (2) Tenax<sup>®</sup> resin tube contents (analyzed together) and the analytical result for the Anasorb 747<sup>®</sup> Tube contents. The calculation is as follows:

(Total  $\mu g$  on the Tenax® Tubes #1 and #2) + (Total  $\mu g$  on the Anasorb 747® Tube) = Total  $\mu g$  on the Method 0031 VOST tube set. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g/\text{set}$ 

When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value included in the tube set total is the default RDL value and the actual value is known to be less than (<) the displayed result.

- The data flags in this column for the VOST Tube Set are the cumulative set of flags contributed by each individual train component included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Tube Set" column when the associated component analytical result is a significant number in comparison to the VOST tube set total. That is, if the VOST Tube Set Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Tube Set Total, but if the VOST Tube Set Total is not affected by a VOST component analytical result, the associated flag is not carried through to the VOST Tube Set Total.
- <sup>4</sup> The VOST Condensate result was obtained by multiplying the sample's corresponding RDL or "hit" by the VOST condensate volume.
- 5. This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - An "E" flag indicates that the result exceeded the upper calibration range. The analytical result is therefore an estimated value.

- A "J" flag indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
- A "B" flag indicates that this compound was found in the associated laboratory method blank. Under these conditions this value is regarded as an estimated value.
- A "Y" flag indicates that this compound is an indistinguishable isomer as a tentatively identified compound (TIC).
- An "N" flag indicates that there is presumptive evidence that this compound is present in the sample based on spectral evidence.
- An "M" flag indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- A "D" flag indicates that this result was obtained by a dilution of the sample. The original analysis yielded an analytical result that exceeded the calibration range.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data.

### **VOST Summary - Run 3 Train Totals** Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-8. HLLWE Run ID: 0031-END-1

Field Sample Name:

Volatile Organic Sampling Train (VOST) Totals
Tenax® and Anasorb 747® Tube Sets (Sets #1, #2, #3, and #4) and the VOST Condensate for Volatile Organic Compounds (VOC) Analysis Sample Description:

		VOS Tube Se (Total με	et #1	VOS Tube S (Total μ	et #2	VOS Tube Se (Total με	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Tota	nsate	VOS Tota (Total	l¹	Power in our
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>						
Target Compound List	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,									.,,				
Acetone	67-64-1	0.70	В	2.6	В	1.4	В	1.1	В	1.4	В	7.2	В	A
Acrylonitrile	107-13-1	< 0.58		< 0.58		< 0.58		< 0.58		< 0.51		< 2.8		N
Benzene	71-43-2	< 0.053		< 0.047		< 0.058		< 0.058		< 0.027		< 0.24		Р
Bromobenzene	108-86-1	< 0.020		< 0.020	***************************************	< 0.020		< 0.020		< 0.036		< 0.12		N
Bromochloromethane	74-97-5	< 0.030		< 0.030	***************************************	< 0.030		< 0.030		< 0.028		< 0.15		N
Bromodichloromethane	75-27-4	< 0.022		< 0.022		< 0.022		< 0.022		< 0.034		< 0.12		N
Bromoform	75-25-2	< 0.038		< 0.038		< 0.038		< 0.038	••••	< 0.026		< 0.18		N
Bromomethane	74-83-9	0.035	J	0.045	J	0.054	J	0.075	J	< 0.020		< 0.23	J	P
2-Butanone	78-93-3	< 0.20		< 0.20	J	< 0.20	J	< 0.20	J	< 0.094		< 0.89	J	P
n-Butylbenzene	104-51-8	< 0.032	***************************************	< 0.032		< 0.032		< 0.032		< 0.026		< 0.15		N
sec-Butylbenzene	135-98-8	< 0.017		< 0.017		< 0.017		< 0.017		< 0.017		< 0.085		N
tert-Butylbenzene	98-06-6	< 0.032	***************************************	< 0.032		< 0.032		< 0.032		< 0.015		< 0.14		N
Carbon disulfide	75-15-0	< 0.35		< 0.17		< 0.39		< 0.27	••••••••••	< 0.011		< 1.2		P
Carbon tetrachloride	56-23-5	< 0.036		< 0.036		< 0.036	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.036		< 0.020		< 0.16		N
Chlorobenzene	108-90-7	< 0.017		< 0.017		< 0.017	J	< 0.017	J	< 0.027		< 0.095	J	P
Chlorodibromomethane	124-48-1	< 0.030		< 0.030		< 0.030		< 0.030		< 0.029		< 0.15		N
Chloroethane	75-00-3	< 0.041	J	< 0.043	J	< 0.047	J	0.049	J	< 0.015		< 0.20	J	P
Chloroform	67-66-3	< 0.096		< 0.090	***************************************	< 0.090		< 0.090		< 0.031		< 0.40		P
Chloromethane	74-87-3	0.52		0.64		0.69		1.0		< 0.011	•••••••••••••••••••••••••••••••••••••••	< 2.9		P
2-Chlorotoluene	95-49-8	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
4-Chlorotoluene	106-43-4	< 0.0094	***************************************	< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
1,2-Dibromo-3-chloropropane	96-12-8	< 0.058		< 0.058	***************************************	< 0.058		< 0.058	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.043		< 0.28		N
1,2-Dibromoethane	106-93-4	< 0.040		< 0.040		< 0.040		< 0.040		< 0.043		< 0.20		N
Dibromomethane	74-95-3	< 0.034	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.034		< 0.034		< 0.034		< 0.030		< 0.17		N
1,2-Dichlorobenzene	95-50-1	< 0.040		< 0.040		< 0.040		< 0.040		< 0.019		< 0.18		N
1,3-Dichlorobenzene	541-73-1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.022		< 0.10		N
1,4-Dichlorobenzene	106-46-7	< 0.028		< 0.028		< 0.028	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.028		< 0.024		< 0.14		N

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### VOST Summary - Run 3 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-8. HLLWE Run ID: 0031-END-1

		VOS Tube Se (Total µg	et #1	VOS Tube S (Total μ	et #2	VOS Tube Se (Total µg	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Tota	nsate	VOS Tota (Total	l <sup>1</sup>	
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>						
Dichlorodifluoromethane	75-71-8	< 0.035	J	< 0.037	J	< 0.034	J	< 0.046	***************************************	< 0.011		< 0.16	J	P
1,1-Dichloroethane	75-34-3	< 0.034		< 0.034		< 0.034		< 0.034		< 0.017		< 0.15		N
1,2-Dichloroethane	107-06-2	< 0.034	J	< 0.020		< 0.16	J	P						
1,1-Dichloroethene	75-35-4	< 0.036	J	< 0.041	J	< 0.039	J	< 0.044	.,,,,,,,,	< 0.016		< 0.18	J	Р
cis-1,2-Dichloroethene	156-59-2	< 0.032	***************************************	< 0.032		< 0.032		< 0.032		< 0.021		< 0.15		N
trans-1,2-Dichloroethene	156-60-5	< 0.038		< 0.038		< 0.038		< 0.038		< 0.013		< 0.16		N
1,2-Dichloropropane	78-87-5	< 0.026		< 0.026		< 0.026		< 0.026	J	< 0.024		< 0.13	J	Р
1,3-Dichloropropane	142-28-9	< 0.038	··········	< 0.038		< 0.038		< 0.038		< 0.022		< 0.17		N
2,2-Dichloropropane	594-20-7	< 0.036		< 0.036		< 0.036		< 0.036		< 0.011		< 0.16		N
1,1-Dichloropropene	563-58-6	< 0.040		< 0.040		< 0.040		< 0.040	,,,,,	< 0.016		< 0.18		N
cis-1,3-Dichloropropene	10061-01-5	< 0.024		< 0.024		< 0.024		< 0.024		< 0.030		< 0.13		N
trans-1,3-Dichloropropene	10061-02-6	< 0.030	***************************************	< 0.030	M	< 0.030	······································	< 0.030		< 0.028		< 0.15	,,,,	N
Ethylbenzene	100-41-4	< 0.018	***************************************	< 0.018		< 0.018	······································	< 0.018		< 0.021		< 0.093		N
Hexachlorobutadiene	87-68-3	< 0.050	***************************************	< 0.050		< 0.050		< 0.050		< 0.025		< 0.22		N
2-Hexanone	591-78-6	< 0.13	***************************************	< 0.13		< 0.13		< 0.13	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.036		< 0.56		N
Isopropylbenzene	98-82-8	< 0.013		< 0.013	***************************************	< 0.013		< 0.013		< 0.018		< 0.070		N
p-Isopropyltoluene	99-87-6	< 0.024		< 0.024		< 0.024		< 0.024		< 0.018		< 0.11		N
Methylene chloride	75-09-2	0.59		0.26		0.42		0.28		0.13	В	1.7	В	A
4-Methyl-2-pentanone	108-10-1	< 0.14		< 0.14		< 0.14	,,,,,,	< 0.14		< 0.030		< 0.59		N
Naphthalene	91-20-3	< 0.050		< 0.050		< 0.050	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.050		< 0.011		< 0.21		N
n-Propylbenzene	103-65-1	< 0.011		< 0.011		< 0.011		< 0.011		< 0.022		< 0.066		N
Styrene	100-42-5	< 0.014		< 0.014		< 0.014		< 0.014	***************************************	< 0.022		< 0.078		N
1,1,1,2-Tetrachloroethane	630-20-6	< 0.019		< 0.019		< 0.019		< 0.019		< 0.024		< 0.10		N
1,1,2,2-Tetrachloroethane	79-34-5	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.22		N
Tetrachloroethene	127-18-4	< 0.032		< 0.032		< 0.032		< 0.032		< 0.021		< 0.15		N
Toluene	108-88-3	0.14		0.051	J	< 0.032	J	< 0.031	J	< 0.028		< 0.28	J	Р
1,2,3-Trichlorobenzene	87-61-6	< 0.050		< 0.050		< 0.050		< 0.050		< 0.011		< 0.21		N
1,2,4-Trichlorobenzene	120-82-1	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.22		N
1,1,1-Trichloroethane	71-55-6	< 0.044		< 0.044	рич	< 0.044		< 0.044		< 0.018		< 0.19		N

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## VOST Summary - Run 3 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-8. HLLWE Run ID: 0031-END-1

	CAS	VOS Tube S (Total μ	et #1	VOS Tube S (Total μ	et #2	VOS Tube So (Total µ)	et #3	VOS Tube S (Total µ	et #4	VO Conde (Tota	nsate	VO Tot (Tota	al <sup>1</sup>	Project
Analyte	Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Specific Flag <sup>6</sup>
1,1,2-Trichloroethane	79-00-5	< 0.036		< 0.036	······································	< 0.036		< 0.036		< 0.022		< 0.17		N
Trichloroethene	79-01-6	< 0.034		< 0.034		< 0.034		< 0.034		< 0.020		< 0.16		N
Trichlorofluoromethane	75-69-4	< 0.036		< 0.036		< 0.036	J	< 0.036		< 0.011		< 0.16	J	P
1,2,3-Trichloropropane	96-18-4	< 0.050		< 0.050		< 0.050		< 0.050		< 0.036		< 0.24		N
1,2,4-Trimethylbenzene	95-63-6	< 0.015		< 0.015		< 0.015		< 0.015		< 0.043		< 0.10		N
1,3,5-Trimethylbenzene	108-67-8	< 0.010		< 0.010		< 0.010		< 0.010	«·····	< 0.019	•	< 0.059		N
Vinyl chloride	75-01-4	< 0.020	J	< 0.025	J	0.023	J	0.039	J	< 0.068	***************************************	< 0.18	J	Р
m-Xylene & p-Xylene	136777-61-2	< 0.10		< 0.10		< 0.10		< 0.10	•••••	< 0.043		< 0.44		N
o-Xylene	95-47-6	< 0.013	J	< 0.013		< 0.013		< 0.013		< 0.026		< 0.078	J	Р
TIC <sup>7</sup>						•••••••••••		••••						
Pentane, 3,3-dimethyl-	562-49-2	0.059										0.059	N,J,M	P
Hexane, 2-methyl-	591-76-4	0.27	***************************************	0.037								0.31	N,J,M	P
Pentane, 2,3-dimethyl-	565-59-3			0.031								0.031	N,J,M	P
Hexane, 3-methyl-	589-34-4	0.65		0.076	•••••••••••••••••••••••••••••••••••••••	0.031						0.76	N,J,M	Р
Cyclohexene	110-83-8	0.036										0.036	N,J,M	Р
Cyclobutane, ethenyl-	2597-49-1			0.029		0.044		0.047				0.12	N,J,M	P
Cyclopentane, 1,2-dimethyl-, t	822-50-4	0.079										0.079	N,J,M	P
Cyclohexane, methyl-	108-87-2	0.20										0.20	N,J,M	P
Hexane, 2,4-dimethyl-	589-43-5	0.16		0.025								0.18	N,J,M	P
Cyclopentane, ethyl-	1640-89-7	0.041										0.041	N,J,M	P
Methane, trichloronitro-	76-06-2							0.36				0.36	N,J,M	P
Benzonitrile	100-47-0			0.074		0.058		0.059				0.19	N,J,M	P
Undecane	1120-21-4	0.21				0.14		0.15				0.50	N,J,M	P
Undecane, 5-methyl-	1632-70-8					0.076		0.10				0.18	N,J,M	Р
Dodecane	112-40-3	8.4		5.7		5.2		6.8				26	N,J,M	P
Tridecane	629-50-5	0.98		0.64		0.58		0.89	DH1D1111111111111111111111111111111111			3.1	N,J,M	Р
Tetradecane	629-59-4	0.40		0.24				0.37				1.0	N,J,M	Р
Hexadecane	544-76-3					0.21						0.21	N,J,M	Р
													1	1

#### Footnotes:

- The Method 0031 VOST Run Total (in Total μg) is the sum of results for the four (4) VOST tube sets and the condensate sample collected during the same sampling run using the following guidelines:
  - When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
  - ♦ When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
  - It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the VOST Total are the cumulative set of flags contributed by each train tube set included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Total" column when the associated component analytical result is a significant number in comparison to the VOST Total. That is, if the VOST Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Total, but if the VOST Total is not affected by a VOST component, the associated flag is not carried through to the VOST Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

The Method 0031 VOST Tube Set (Total μg/Set) result consists of the sum of the analytical results for the two (2) Tenax<sup>®</sup> resin tube contents (analyzed together) and the analytical result for the Anasorb 747<sup>®</sup> Tube contents. The calculation is as follows:

(Total  $\mu g$  on the Tenax® Tubes #1 and #2) + (Total  $\mu g$  on the Anasorb 747® Tube) = Total  $\mu g$  on the Method 0031 VOST tube set. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g/\text{set}$ 

When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value included in the tube set total is the default RDL value and the actual value is known to be less than (<) the displayed result.

- The data flags in this column for the VOST Tube Set are the cumulative set of flags contributed by each individual train component included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Tube Set" column when the associated component analytical result is a significant number in comparison to the VOST tube set total. That is, if the VOST Tube Set Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Tube Set Total, but if the VOST Tube Set Total is not affected by a VOST component analytical result, the associated flag is not carried through to the VOST Tube Set Total.
- <sup>4</sup> The VOST Condensate result was obtained by multiplying the sample's corresponding RDL or "hit" by the VOST condensate volume.
- <sup>5</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - An "E" flag indicates that the result exceeded the upper calibration range. The analytical result is therefore an estimated value.

- A "J" flag indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
- A "B" flag indicates that this compound was found in the associated laboratory method blank. Under these conditions this value is regarded as an estimated value.
- ♦ A "Y" flag indicates that this compound is an indistinguishable isomer as a tentatively identified compound (TIC).
- An "N" flag indicates that there is presumptive evidence that this compound is present in the sample based on spectral evidence.
- An "M" flag indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- ♦ A "D" flag indicates that this result was obtained by a dilution of the sample. The original analysis yielded an analytical result that exceeded the calibration range.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- 7. The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data.

### VOST Summary - Run 2 Train Totals Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-9. HLLWE Run ID: 0031-STRT-2

Field Sample Name: Volatile Organic Sampling Train (VOST) Totals

Sample Description: Tenax® and Anasorb 747® Tube Sets (Sets #1, #2, #3, and #4) and the VOST Condensate for Volatile Organic Compounds (VOC) Analysis

		VOS Tube S (Total μ	et#1	VOS Tube S (Total μ	et #2	VOS Tube Se (Total μ	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Total	nsate	VOS Tota (Total	al¹	
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
Target Compound List													••••••	
Acetone	67-64-1	1.2	В	0.81	J,B	1.2	В	< 0.26	J,B	1.5	В	5.0	J,B	A
Acrylonitrile	107-13-1	< 0.58		< 0.58		< 0.58		< 0.58		< 0.51		< 2.8		N
Benzene	71-43-2	< 0.063		< 0.053		< 0.040	J	< 0.038	J	< 0.027		< 0.22	J	Р
Bromobenzene	108-86-1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.036		< 0.12		N
Bromochloromethane	74-97-5	< 0.030		< 0.030		< 0.030		< 0.030		< 0.028		< 0.15		N
Bromodichloromethane	75-27-4	< 0.022		< 0.022		< 0.022		< 0.022		< 0.034		< 0.12		N
Bromoform	75-25-2	< 0.038		< 0.038		< 0.038		< 0.038		< 0.026		< 0.18		N
Bromomethane	74-83-9	< 0.030	J	< 0.030	J	< 0.033	J	< 0.039	J	< 0.020		< 0.15	J	P
2-Butanone	78-93-3	< 0.20		< 0.20	••••••	< 0.20	J	< 0.20		< 0.094		< 0.89	J	Р
n-Butylbenzene	104-51-8	< 0.032		< 0.032		< 0.032		< 0.032		< 0.026		< 0.15		N
sec-Butylbenzene	135-98-8	< 0.017		< 0.017		< 0.017		< 0.017		< 0.017		< 0.085		N
tert-Butylbenzene	98-06-6	< 0.032		< 0.032	***************************************	< 0.032		< 0.032		< 0.015		< 0.14		N
Carbon disulfide	75-15-0	< 0.46		< 0.31		< 0.29		< 0.26		< 0.011		< 1.3		P
Carbon tetrachloride	56-23-5	< 0.036		< 0.036		< 0.036		< 0.036		< 0.020		< 0.16		N
Chlorobenzene	108-90-7	< 0.017		< 0.017		< 0.017		< 0.017		< 0.027		< 0.095		N
Chlorodibromomethane	124-48-1	< 0.030		< 0.030		< 0.030		< 0.030		< 0.029		< 0.15		N
Chloroethane	75-00-3	< 0.036		< 0.036		< 0.042	J	< 0.038	J	< 0.015		< 0.17	J	Р
Chloroform	67-66-3	< 0.036		< 0.036		< 0.094		< 0.13		< 0.031		< 0.33		Р
Chloromethane	74-87-3	< 0.061	J	0.16	J	0.47		0.48		< 0.011		< 1.2	J	P
2-Chlorotoluene	95 <b>-</b> 49-8	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
4-Chlorotoluene	106-43-4	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
1,2-Dibromo-3-chloropropane	96-12-8	< 0.058		< 0.058		< 0.058		< 0.058	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.043		< 0.28		N
1,2-Dibromoethane	106-93-4	< 0.040		< 0.040		< 0.040		< 0.040		< 0.043		< 0.20		N
Dibromomethane	74-95-3	< 0.034		< 0.034		< 0.034		< 0.034		< 0.030		< 0.17		N
1,2-Dichlorobenzene	95-50-1	< 0.040		< 0.040		< 0.040		< 0.040		< 0.019		< 0.18		N
1,3-Dichlorobenzene	541-73-1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.022		< 0.10		N

### VOST Summary - Run 2 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-9. HLLWE Run ID: 0031-STRT-2

		VOST Tube Set #1 (Total µg/Set)	VOST Tube Set #2 (Total µg/Set)	VOST Tube Set #3 (Total µg/Set)	VOST Tube Set #4 (Total µg/Set)	VOST Condensate (Total µg)	VOST Total¹ (Total μg)	
Analyte	CAS Registry Number	Risk Result <sup>2</sup> Flag <sup>3</sup>	Risk Result <sup>4</sup> Flag <sup>5</sup>	Total Flag	Project Specific Flag <sup>6</sup>			
1,4-Dichlorobenzene	106-46-7	< 0.028	< 0.028	< 0.028	< 0.028	< 0.024	< 0.14	N
Dichlorodifluoromethane	75-71-8	< 0.047	< 0.054	< 0.048	< 0.039	< 0.011	< 0.20	P
1,1-Dichloroethane	75-34-3	< 0.034	< 0.034	< 0.034	< 0.034	< 0.017	< 0.15	N
1,2-Dichloroethane	107-06-2	< 0.034	< 0.034	< 0.034	< 0.034	< 0.020	< 0.16	N
1,1-Dichloroethene	75-35-4	< 0.036	< 0.036	< 0.036 J	< 0.036 J	< 0.016	< 0.16 J	Р
cis-1,2-Dichloroethene	156-59-2	< 0.032	< 0.032	< 0.032	< 0.032	< 0.021	< 0.15	N
trans-1,2-Dichloroethene	156-60-5	< 0.038	< 0.038	< 0.038	< 0.038	< 0.013	< 0.16	N
1,2-Dichloropropane	78-87-5	< 0.026	< 0.026	< 0.026	< 0.026	< 0.024	< 0.13	N
1,3-Dichloropropane	142-28-9	< 0.038	< 0.038	< 0.038	< 0.038	< 0.022	< 0.17	N
2,2-Dichloropropane	594-20-7	< 0.036	< 0.036	< 0.036	< 0.036	< 0.011	< 0.16	N
I,l-Dichloropropene	563-58-6	< 0.040	< 0.040	< 0.040	< 0.040	< 0.016	< 0.18	N
cis-1,3-Dichloropropene	10061-01-5	< 0.024	< 0.024	< 0.024	< 0.024	< 0.030	< 0.13	N
trans-1,3-Dichloropropene	10061-02-6	< 0.030	< 0.030	< 0.030	< 0.030	< 0.028	< 0.15	N
Ethylbenzene	100-41-4	< 0.018	< 0.018	< 0.018	< 0.018	< 0.021	< 0.093	N
Hexachlorobutadiene	87-68-3	< 0.050	< 0.050	< 0.050	< 0.050	< 0.025	< 0.22	N
2-Hexanone	591-78-6	< 0.13	< 0.13	< 0.13	< 0.13	< 0.036	< 0.56	N
Isopropylbenzene	98-82-8	< 0.013	< 0.013	< 0.013	< 0.013	< 0.018	< 0.070	N
p-Isopropyltoluene	99-87-6	< 0.024	< 0.024	< 0.024	< 0.024	< 0.018	< 0.11	N
Methylene chloride	75-09-2	0.55 B	0.15 B	0.13 B	0.12 B	0.13 B	1.1 B	А
4-Methyl-2-pentanone	108-10-1	< 0.14	< 0.14	< 0.14	< 0.14	< 0.030	< 0.59	N
Naphthalene	91-20-3	< 0.050	< 0.050	< 0.050	< 0.050	< 0.011	< 0.21	N
n-Propylbenzene	103-65-1	< 0.011	< 0.011	< 0.011	< 0.011	< 0.022	< 0.066	N
Styrene	100-42-5	< 0.014	< 0.014	< 0.014	< 0.014	< 0.022	< 0.078	N
1,1,1,2-Tetrachloroethane	630-20-6	< 0.019	< 0.019	< 0.019	< 0.019	< 0.024	< 0.10	N
1,1,2,2-Tetrachloroethane	79-34-5	< 0.050	< 0.050	< 0.050	< 0.050	< 0.025	< 0.22	N
Tetrachloroethene	127-18-4	< 0.032	< 0.032	< 0.032	< 0.032	< 0.021	< 0.15	N
Toluene	108-88-3	< 0.027 J	0.17 J	0.031 J	· 0.045 J	< 0.028	< 0.30 J	Р
1,2,3-Trichlorobenzene	87-61-6	< 0.050	< 0.050	< 0.050	< 0.050	< 0.011	< 0.21	N
1,2,4-Trichlorobenzene	120-82-1	< 0.050	< 0.050	< 0.050	< 0.050	< 0.025	< 0.22	N

## VOST Summary - Run 2 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-9. HLLWE Run ID: 0031-STRT-2

		VOST Tube Se (Total μg	t #1	VOS Tube S (Total μ	et #2	VOS Tube So (Total με	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Tota	nsate	VOS Tota (Total	al <sup>i</sup>	
Analyte	CAS Registry Number	Risk Result²	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
1,1,1-Trichloroethane	71-55-6	< 0.044		< 0.044		< 0.044		< 0.044		< 0.018	······································	< 0.19		N
1,1,2-Trichloroethane	79-00-5	< 0.036	***************************************	< 0.036		< 0.036		< 0.036		< 0.022		< 0.17		N
Trichloroethene	79-01-6	< 0.034		< 0.034		< 0.034		< 0.034		< 0.020		< 0.16		N
Trichlorofluoromethane	75-69-4	< 0.036	J	< 0.036	J	< 0.036	J	< 0.036		< 0.011		< 0.16	J	P
1,2,3-Trichloropropane	96-18-4	< 0.050		< 0.050		< 0.050		< 0.050		< 0.036		< 0.24		N
1,2,4-Trimethylbenzene	95-63-6	< 0.015		< 0.015		< 0.015		< 0.015		< 0.043		< 0.10		N
1,3,5-Trimethylbenzene	108-67-8	< 0.010		< 0.010	***************************************	< 0.010		< 0.010	***************************************	< 0.019		< 0.059		N
Vinyl chloride	75-01-4	< 0.013		< 0.013		< 0.018	J	< 0.016	J	< 0.068		< 0.13	J	P
m-Xylene & p-Xylene	136777-61-2	< 0.10		< 0.10	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.10		< 0.10		< 0.043		< 0.44		N
o-Xylene	95-47-6	< 0.013		< 0.013		< 0.013		< 0.013		< 0.026		< 0.078		N

### VOST Summary - Run 2 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-9. HLLWE Run ID: 0031-STRT-2

		VOST Tube Set #1 (Total μg/Set)	VOST Tube Set #2 (Total μg/Set)	VOST Tube Set #3 (Total μg/Set)	VOST Tube Set #4 (Total μg/Set)	VOST Condensate (Total μg)	VOST Total¹ (Total μg)	
Analyte	CAS Registry Number	Risk Result <sup>2</sup> Flag <sup>3</sup>	Risk Result <sup>4</sup> Flag <sup>5</sup>	Total Flag	Project Specific Flag <sup>6</sup>			
TICs <sup>7</sup>				***************************************	!			
Hexane, 2-methyl-	591-76-4		0.17	0.059	0.072		0.30 N,J,M	Р
Pentane, 2,3-dimethyl-	565-59-3			0.057	0.056		0.11 N,J,M	P
Hexane, 3-methyl-	589-34-4			0.12	0.15		0.27 N,J,M	Р
Pentane, 3-ethyl-	617-78-7			0.034			0.034 N,J,M	Р
Cyclohexene	110-83-8	0.030	0.042	0.036	0.073		0.18 N,J,M	Р
Cyclopentane, 1,2-dimethyl-	2452-99-5		0.053				0.053 N,J,M	Р
Cyclohexane, methyl-	108-87-2		0.13	0.032	0.037		0.20 N,J,M	P
Hexane, 2,4-dimethyl-	589-43-5		0.13	0.037	0.037		0.20 N,J,M	P
Cyclopentane, ethyl-	1640-89-7		0.036				0.036 N,J,M	P
Benzonitrile	100-47-0	0.038	0.034				0.072 N,J,M	P
Tridecane	629-50-5	0.082					0.082 N,J,M	Р
Undecane	1120-21-4		0.065	0.10	0.080		0.24 N,J,M	Р
Decane, 2,2,5-trimethyl-	62237-96-1				0.060		0.060 N,J,M	Р
Undecane, 5-methyl-	1632-70-8	0.052	0.051	0.11			0.21 N,J,M	P
Dodecane	112-40-3	3.4	3.4	8.1	6.8		22 N,J,M	Р
Dodecane, 6-methyl-	6044-71-9			0.052			0.052 N,J,M	Р
Undecane, 2,6-dimethyl-	17301-23-4				0.046		0.046 N,J,M	Р
Tridecane	629-50-5	0.34	0.29	0.84	0.93		2.4 N,J,M	P
Tetradecane	629-59-4	0.17	0.16	0.28	0.37		0.98 N,J,M	Р

#### Footnotes:

- The Method 0031 VOST Run Total (in Total μg) is the sum of results for the four (4) VOST tube sets and the condensate sample collected during the same sampling run using the following guidelines:
  - When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
  - When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
  - It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the VOST Total are the cumulative set of flags contributed by each train tube set included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Total" column when the associated component analytical result is a significant number in comparison to the VOST Total. That is, if the VOST Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Total, but if the VOST Total is not affected by a VOST component, the associated flag is not carried through to the VOST Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

The Method 0031 VOST Tube Set (Total μg/Set) result consists of the sum of the analytical results for the two (2) Tenax<sup>®</sup> resin tube contents (analyzed together) and the analytical result for the Anasorb 747<sup>®</sup> Tube contents. The calculation is as follows:

(Total  $\mu g$  on the Tenax® Tubes #1 and #2) + (Total  $\mu g$  on the Anasorb 747® Tube) = Total  $\mu g$  on the Method 0031 VOST tube set. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g/\text{set}$ 

When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value included in the tube set total is the default RDL value and the actual value is known to be less than (<) the displayed result.

- The data flags in this column for the VOST Tube Set are the cumulative set of flags contributed by each individual train component included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Tube Set" column when the associated component analytical result is a significant number in comparison to the VOST tube set total. That is, if the VOST Tube Set Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Tube Set Total, but if the VOST Tube Set Total is not affected by a VOST component analytical result, the associated flag is not carried through to the VOST Tube Set Total.
- <sup>4</sup> The VOST Condensate result was obtained by multiplying the sample's corresponding RDL or "hit" by the VOST condensate volume.
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - An "E" flag indicates that the result exceeded the upper calibration range. The analytical result is therefore an estimated value.

- A "J" flag indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
- A "B" flag indicates that this compound was found in the associated laboratory method blank. Under these conditions this value is regarded as an estimated value.
- A "Y" flag indicates that this compound is an indistinguishable isomer as a tentatively identified compound (TIC).
- An "N" flag indicates that there is presumptive evidence that this compound is present in the sample based on spectral evidence.
- An "M" flag indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- A "D" flag indicates that this result was obtained by a dilution of the sample. The original analysis yielded an analytical result that exceeded the calibration range.
- 6. Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- 7. The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data.

### **VOST Summary - Run 4 Train Totals** Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-10. HLLWE Run ID: 0031-END-2

Field Sample Name:

Volatile Organic Sampling Train (VOST) Totals
Tenax® and Anasorb 747® Tube Sets (Sets #1, #2, #3, and #4) and the VOST Condensate for Volatile Organic Compounds (VOC) Analysis Sample Description:

	010	VOS Tube Se (Total μ	et #1	VOS Tube S (Total μ	et #2	VOS Tube Se (Total µg	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Total	nsate	VOS Tota (Total	ıl¹	Project
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Specific Flag <sup>6</sup>
Target Compound List													,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
Acetone	67-64-1	< 0.46	J,B	0.59	В	1.3	В	1.4	В	0.081	J,B	< 3.8	J,B	A
Acrylonitrile	107-13-1	< 0.58		< 0.58		< 0.58		< 0.58		< 0.51	***************************************	< 2.8		N
Benzene	71-43-2	< 0.052		< 0.034		< 0.044		< 0.047		< 0.027		< 0.20		P
Bromobenzene	108-86-1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.036		< 0.12		N
Bromochloromethane	74-97-5	< 0.030	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.030		< 0.030		< 0.030		< 0.028		< 0.15		N
Bromodichloromethane	75-27-4	< 0.022		< 0.022		< 0.022		< 0.022		< 0.033		< 0.12		N
Bromoform	75-25-2	< 0.038	,,	< 0.038		< 0.038	***************************************	< 0.038		< 0.025		< 0.18		N
Bromomethane	74-83-9	< 0.030	J	< 0.055	J	< 0.041	J	< 0.049	J	< 0.020		< 0.20	J	Р
2-Butanone	78-93-3	< 0.20	***************************************	< 0.20		< 0.20		< 0.20		< 0.093		< 0.89		N
n-Butylbenzene	104-51-8	< 0.032		< 0.032		< 0.032		< 0.032		< 0.025		< 0.15		N
sec-Butylbenzene	135-98-8	< 0.017		< 0.017		< 0.017	***************************************	< 0.017		< 0.017		< 0.085		N
tert-Butylbenzene	98-06-6	< 0.032	***************************************	< 0.032		< 0.032		< 0.032		< 0.014		< 0.14		N
Carbon disulfide	75-15-0	0.22		< 0.036		< 0.18		< 0.19		< 0.011		< 0.64		Р
Carbon tetrachloride	56-23-5	< 0.036	J	< 0.036		< 0.036	••••	< 0.036	J	< 0.020		< 0.16	J	P
Chlorobenzene	108-90-7	< 0.017	J	< 0.017		< 0.017	J	< 0.017	J	< 0.027		< 0.095	J	P
Chlorodibromomethane	124-48-1	< 0.030		< 0.030		< 0.030		< 0.030		< 0.029		< 0.15		N
Chloroethane	75-00-3	< 0.036	J	< 0.036	J	< 0.036	J	< 0.036	J	< 0.014		< 0.16	J	P
Chloroform	67-66-3	< 0.15		< 0.036		< 0.087		< 0.087		< 0.031		< 0.39		Р
Chloromethane	74-87-3	0.46		< 0.70		0.65	J	0.70		< 0.011		< 2.5	J	P
2-Chlorotoluene	95-49-8	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
4-Chlorotoluene	106-43-4	< 0.0094		< 0.0094		< 0.0094		< 0.0094		< 0.021		< 0.059		N
1,2-Dibromo-3-chloropropane	96-12-8	< 0.058		< 0.058		< 0.058		< 0.058		< 0.042		< 0.27		N
1,2-Dibromoethane	106-93-4	< 0.040		< 0.040		< 0.040		< 0.040		< 0.042		< 0.20		N
Dibromomethane	74-95-3	< 0.034		< 0.034	••••	< 0.034		< 0.034	.,	< 0.030		< 0.17		N
1,2-Dichlorobenzene	95-50-1	< 0.040		< 0.040		< 0.040		< 0.040		< 0.019		< 0.18		N
1,3-Dichlorobenzene	541-73 <b>-</b> 1	< 0.020		< 0.020		< 0.020		< 0.020		< 0.022		< 0.10		N
1,4-Dichlorobenzene	106-46-7	< 0.028		< 0.028	***************************************	< 0.028		< 0.028		< 0.023		< 0.14		N
	***************************************	A												

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## VOST Summary - Run 4 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-10. HLLWE Run ID: 0031-END-2

		VOS Tube Se (Total µ	et #1	VOS Tube S (Total µ	et #2	VOS Tube Se (Total με	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Total	nsate	VOS Tota (Total	d <sup>1</sup>	
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
Dichlorodifluoromethane	75-71-8	< 0.032	J	< 0.041		< 0.039		< 0.039		< 0.011		< 0.16	J	P
1,1-Dichloroethane	75-34-3	< 0.034		< 0.034		< 0.034	••••	< 0.034		< 0.017		< 0.15		N
1,2-Dichloroethane	107-06-2	< 0.034		< 0.034		< 0.034		< 0.034		< 0.020		< 0.16		N
I,I-Dichloroethene	75-35-4	< 0.036	J	< 0.036		< 0.036	J	< 0.036	J	< 0.016		< 0.16	J	P
cis-1,2-Dichloroethene	156-59-2	< 0.032		< 0.032		< 0.032		< 0.032		< 0.021		< 0.15		N
trans-1,2-Dichloroethene	156-60-5	< 0.038		< 0.038	haner	< 0.038		< 0.038		< 0.013		< 0.17		N
1,2-Dichloropropane	78-87-5	< 0.026		< 0.026		< 0.026		< 0.026		< 0.023		< 0.13		N
1,3-Dichloropropane	142-28-9	< 0.038		< 0.038		< 0.038		< 0.038		< 0.022		< 0.17		N
2,2-Dichloropropane	594-20-7	< 0.036	·,····································	< 0.036		< 0.036		< 0.036		< 0.011		< 0.16		N
1,1-Dichloropropene	563-58-6	< 0.040		< 0.040		< 0.040		< 0.040		< 0.016		< 0.18		N
cis-1,3-Dichloropropene	10061-01-5	< 0.024	***************************************	< 0.024		< 0.024		< 0.024		< 0.030		< 0.13		N
trans-1,3-Dichloropropene	10061-02-6	< 0.030		< 0.030		< 0.030		< 0.030		< 0.028		< 0.15		N
Ethylbenzene	100-41-4	< 0.018		< 0.018		< 0.018		< 0.018		< 0.021		< 0.093		N
Hexachlorobutadiene	87-68-3	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.23		N
2-Hexanone	591-78-6	< 0.13	***************************************	< 0.13		< 0.13		< 0.13	•	< 0.036		< 0.56		N
Isopropylbenzene	98-82-8	< 0.013		< 0.013	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.013		< 0.013	***************************************	< 0.018		< 0.070		N
p-Isopropyltoluene	99-87-6	< 0.024		< 0.024		< 0.024		< 0.024		< 0.018		< 0.11	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	N
Methylene chloride	75-09-2	0.13	В	0.071	В	0.10	В	0.11	В	< 0.019	J,B	< 0.43	J,B	A
4-Methyl-2-pentanone	108-10-1	< 0.14		< 0.14		< 0.14		< 0.14		< 0.030		< 0.59		N
Naphthalene	91-20-3	< 0.050		< 0.050		< 0.050		< 0.050		< 0.011		< 0.21		N
n-Propylbenzene	103-65-1	< 0.011		< 0.011		< 0.011		< 0.011		< 0.022		< 0.066		N
Styrene	100-42-5	< 0.014		< 0.014		< 0.014		< 0.014		< 0.022		< 0.078		N
1,1,1,2-Tetrachloroethane	630-20-6	< 0.019		< 0.019		< 0.019		< 0.019		< 0.023		< 0.099		N
1,1,2,2-Tetrachloroethane	79-34-5	< 0.050		< 0.050		< 0.050		< 0.050	***************************************	< 0.025		< 0.23		N
Tetrachloroethene	127-18-4	< 0.032		< 0.032		< 0.032		< 0.032		< 0.021		< 0.15		N
Toluene	108-88-3	< 0.048		< 0.046		< 0.028	J	0.026	J	< 0.028	101,000,000,000	< 0.18	J	Р
1,2,3-Trichlorobenzene	87-61-6	< 0.050		< 0.050	,,	< 0.050		< 0.050		< 0.011		< 0.21		N
1,2,4-Trichlorobenzene	120-82-1	< 0.050		< 0.050		< 0.050		< 0.050		< 0.025		< 0.23		N
1.1,1-Trichloroethane	71-55-6	< 0.044	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.044		< 0.044		< 0.044		< 0.018	,	< 0.19		N

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## VOST Summary - Run 4 Train Totals (Continued) Method 0031 Volatile Organic Compounds Analytical Results Summary Table A-10. HLLWE Run ID: 0031-END-2

		VOS Tube Se (Total με	et #1	VOS Tube Se (Total μ	et #2	VOS Tube Se (Total με	et #3	VOS Tube S (Total μ	et #4	VOS Conde (Total	nsate	VOS Tota (Tota	ai¹	
Analyte	CAS Registry Number	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>2</sup>	Flag <sup>3</sup>	Risk Result <sup>4</sup>	Flag <sup>5</sup>	Total	Flag	Project Specific Flag <sup>6</sup>
1,1,2-Trichloroethane	79-00-5	< 0.036		< 0.036		< 0.036		< 0.036		< 0.022		< 0.17		N
Trichloroethene	79-01-6	< 0.034		< 0.034		< 0.034		< 0.034	***************************************	< 0.020		< 0.16		N
Trichlorofluoromethane	75-69-4	< 0.036	J	< 0.036		< 0.036	J	< 0.036	J	< 0.011	ed miles administer	< 0.16	J	Р
1,2,3-Trichloropropane	96-18-4	< 0.050		< 0.050		< 0.050	,	< 0.050		< 0.036		< 0.24		N
1,2,4-Trimethylbenzene	95-63-6	< 0.015		< 0.015		< 0.015		< 0.015	***************************************	< 0.042		< 0.10		N
1,3,5-Trimethylbenzene	108-67-8	< 0.010		< 0.010		< 0.010		< 0.010	•••••	< 0.019		< 0.059		N
Vinyl chloride	75-01-4	< 0.020	J	< 0.026	J	< 0.026	J	< 0.027	J	< 0.068		< 0.17	J	Р
m-Xylene & p-Xylene	136777-61-2	< 0.10	.,,	< 0.10		< 0.10		< 0.10	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	< 0.042		< 0.44		N
o-Xylene	95-47-6	< 0.013	J	< 0.013		< 0.013		< 0.013		< 0.025		< 0.077	J	Р
TICs <sup>7</sup>		•••••••••••••••••••••••••••••••••••••••							,					
Hexane, 2-methyl-	591-76-4	0.047	,	0.10		0.052		0.040				0.24	N,J,M	P
Pentane, 2,3-dimethyl-	565-59-3			0.10				0.037				0.14	N,J,M	P
Hexane, 3-methyl-	589-34-4			0.26				0.081				0.34	N,J,M	Р
Pentane, 3-ethyl-	617-78-7					0.032					***************************************	0.032	N,J,M	P
Cyclohexene	110-83-8	0.026										0.026	N,J,M	Р
Cyclopentane, 1,2-dimethyl-, t	822-50-4			0.030								0.030	N,J,M	P
Cyclohexane, methyl-	108-87-2	0.027		0.065		0.031						0.12	N,J,M	P
Hexane, 2,4-dimethyl-	589-43-5			0.066		0.028						0.094	N,J,M	P
Benzonitrile	100-47-0							0.047				0.047	N,J,M	Р
Undecane	1120-21-4	0.014				0.054		0.049				0.12	N,J,M	Р
Undecane, 5-methyl-	1632-70-8	0.13				0.047						0.18	N,J,M	Р
Dodecane	112-40-3	9.9		0.070		3.8		3.5				17	N,J,M	P
Undecane, 2,6-dimethyl-	17301-23-4	0.083										0.083	N,J,M	Р
Tridecane	629-50-5	1.9		0.046		0.58		0.53				3.1	N,J,M	P
Tetradecane	629-59-4	0.58		0.096		0.35	***************************************	0.36				1.4	N,J,M	Р

#### Footnotes:

- The Method 0031 VOST Run Total (in Total μg) is the sum of results for the four (4) VOST tube sets and the condensate sample collected during the same sampling run using the following guidelines:
  - When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
  - When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "J" flag is carried through the calculation to the train total.
  - When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
  - It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the VOST Total are the cumulative set of flags contributed by each train tube set included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Total" column when the associated component analytical result is a significant number in comparison to the VOST Total. That is, if the VOST Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Total, but if the VOST Total is not affected by a VOST component, the associated flag is not carried through to the VOST Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

The Method 0031 VOST Tube Set (Total  $\mu$ g/Set) result consists of the sum of the analytical results for the two (2) Tenax<sup>®</sup> resin tube contents (analyzed together) and the analytical result for the Anasorb 747<sup>®</sup> Tube contents. The calculation is as follows:

(Total  $\mu g$  on the Tenax<sup>®</sup> Tubes #1 and #2) + (Total  $\mu g$  on the Anasorb 747<sup>®</sup> Tube) = Total  $\mu g$  on the Method 0031 VOST tube set. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g/\text{set}$ 

When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value included in the tube set total is the default RDL value and the actual value is known to be less than (<) the displayed result.

- The data flags in this column for the VOST Tube Set are the cumulative set of flags contributed by each individual train component included as part of the VOST total. A flag attached to a VOST component is carried through to the "VOST Tube Set" column when the associated component analytical result is a significant number in comparison to the VOST tube set total. That is, if the VOST Tube Set Total is affected by a VOST component analytical result, the associated flag is carried through to the VOST Tube Set Total, but if the VOST Tube Set Total is not affected by a VOST component analytical result, the associated flag is not carried through to the VOST Tube Set Total.
- <sup>4</sup> The VOST Condensate result was obtained by multiplying the sample's corresponding RDL or "hit" by the VOST condensate volume.
- <sup>5</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - ♦ A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - An "E" flag indicates that the result exceeded the upper calibration range. The analytical result is therefore an estimated value.

- A "J" flag indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
- A "B" flag indicates that this compound was found in the associated laboratory method blank. Under these conditions this value is regarded as an estimated value.
- A "Y" flag indicates that this compound is an indistinguishable isomer as a tentatively identified compound (TIC).
- An "N" flag indicates that there is presumptive evidence that this compound is present in the sample based on spectral evidence.
- An "M" flag indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
- A "D" flag indicates that this result was obtained by a dilution of the sample. The original analysis yielded an analytical result that exceeded the calibration range.
- <sup>6</sup> Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.
- 7. The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data.

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### BECHTEL BWXT IDAHO, LLC (BBWI) INTEC HLLWE Effluent Gas Emissions Inventory Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### VOST Analytical Results Summary Table A-11. Run 2, VOST Tenax® Tube Pair Field Blank

Field Sample Name:

Volatile Organic Sampling Train (VOST)

Sample Description:

VOST Tenax® Tube Pair Field Blank for Volatile Organic Compounds Analysis

Field Sample Number(s):

A-3392

STL Sample Number(s).

H1F250144-019

	CAS Registry			Tenax® T Field (μg/Sa	Blank		
Analyte	Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL⁴	Risk Result <sup>5</sup>	Flag <sup>6</sup>
Acetone	67-64-1	0.027	0.024	0.063	0.10	< 0.063	J,B
Acrylonitrile	107-13-1	ND	0.11	0.29	0.50	< 0.29	
Benzene	71-43-2	ND	0.0064	0.017	0.025	< 0.017	
Bromobenzene	108-86-1	ND	0.0039	0.010	0.025	< 0.010	
Bromochloromethane	74-97-5	ND	0.0056	0.015	0.025	< 0.015	
Bromodichloromethane	75-27-4	ND	0.0042	0.011	0.025	< 0.011	
Bromoform	75-25-2	ND	0.0074	0.019	0.025	< 0.019	
Bromomethane	74-83-9	ND	0.0059	0.015	0.050	< 0.015	,
2-Butanone	78-93-3	ND	0.038	0.10	0.10	< 0.10	
n-Butylbenzene	104-51-8	ND	0.0061	0.016	0.025	< 0.016	
sec-Butylbenzene	135-98-8	ND	0.0032	0.0084	0.025	< 0.0084	
tert-Butylbenzene	98-06-6	ND	0.0062	0.016	0.025	< 0.016	
Carbon disulfide	75-15-0	ND	0.0070	0.018	0.025	< 0.018	
Carbon tetrachloride	56-23-5	ND	0.0069	0.018	0.025	< 0.018	
Chlorobenzene	108-90-7	ND	0.0032	0.0084	0.025	< 0.0084	
Chlorodibromomethane	124-48-1	ND	0.0056	0.015	0.025	< 0.015	
Chloroethane	75-00 <b>-</b> 3	ND	0.0068	0.018	0.050	< 0.018	
Chloroform	67-66-3	ND	0.0070	0.018	0.025	< 0.018	
Chloromethane	74-87-3	ND	0.0048	0.013	0.050	< 0.013	
2-Chlorotoluene	95-49-8	ND	0.0018	0.0047	0.025	< 0.0047	
4-Chlorotoluene	106-43-4	ND	0.0018	0.0047	0.025	< 0.0047	
1,2-Dibromo-3-chloropropane	96-12-8	ND	0.011	0.029	0.050	< 0.029	
1,2-Dibromoethane	106-93-4	ND	0.0075	0.020	0.025	< 0.020	
Dibromomethane	74-95-3	ND	0.0064	0.017	0.025	< 0.017	
1,2-Dichlorobenzene	95-50-1	ND	0.0077	0.020	0.025	< 0.020	
1,3-Dichlorobenzene	541-73-1	ND	0.0038	0.010	0.025	< 0.010	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
1,4-Dichlorobenzene	106-46-7	ND	0.0055	0.014	0.025	< 0.014	
Dichlorodifluoromethane	75-71-8	ND	0.0051	0.013	0.025	< 0.013	

VOST Analytical Results Summary (Continued)
Table A-11. Run 2, VOST Tenax® Tube Pair Field Blank

, 400	CAS Registry			Tenax® T Field (µg/Sa			
Analyte	Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result <sup>5</sup>	Flag
1,1-Dichloroethane	75-34-3	ND	0.0064	0.017	0.025	< 0.017	
1,2-Dichloroethane	107-06-2	ND	0.0066	0.017	0.025	< 0.017	
1.1-Dichloroethene	75-35-4	ND	0.0067	0.018	0.025	< 0.018	
cis-1,2-Dichloroethene	156-59-2	ND	0.0062	0.016	0.025	< 0.016	·····
trans-1,2-Dichloroethene	156-60-5	ND	0.0074	0.019	0.025	< 0.019	
1,2-Dichloropropane	78-87-5	ND	0.0049	0.013	0.025	< 0.013	
1,3-Dichloropropane	142-28-9	ND	0.0073	0.019	0.025	< 0.019	
2,2-Dichloropropane	594-20-7	ND	0.0070	0.018	0.025	< 0.018	
1,1-Dichloropropene	563-58-6	ND	0.0077	0.020	0.025	< 0.020	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
cis-1,3-Dichloropropene	10061-01-5	ND	0.0046	0.012	0.025	< 0.012	
trans-1,3-Dichloropropene	10061-02-6	ND	0.0059	0.015	0.025	< 0.015	
Ethylbenzene	100-41-4	ND	0.0035	0.0092	0.025	< 0.0092	
Hexachlorobutadiene	87-68-3	ND	0.012	0.031	0.025	< 0.025	***************************************
2-Hexanone	591-78-6	ND	0.024	0.063	0.10	< 0.063	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
Isopropylbenzene	98-82-8	ND	0.0024	0.0063	0.025	< 0.0063	
p-lsopropyltoluene	99-87-6	ND	0.0044	0.012	0.025	< 0.012	
Methylene chloride	75-09-2	0.025	0.016	0.042	0.025	0.025	В
4-Methyl-2-pentanone	108-10-1	ND	0.027	0.071	0.10	< 0.071	
Naphthalene	91-20-3	ND	0.014	0.037	0.025	< 0.025	,,,
n-Propylbenzene	103-65-1	ND	0.0021	0.0055	0.025	< 0.0055	
Styrene	100-42-5	ND	0.0026	0.0068	0.025	< 0.0068	
1,1,1,2-Tetrachloroethane	630-20-6	ND	0.0037	0.0097	0.025	< 0.0097	
1,1,2,2-Tetrachloroethane	79-34-5	ND	0.0097	0.025	0.025	< 0.025	
Tetrachloroethene	127-18-4	ND	0.0062	0.016	0.025	< 0.016	
Toluene	108-88-3	ND	0.0025	0.0066	0.025	< 0.0066	
1,2,3-Trichlorobenzene	87-61-6	ND	0.014	0.037	0.025	< 0.025	
1,2,4-Trichlorobenzene	120-82-1	ND	0.013	0.034	0.025	< 0.025	
1,1,1-Trichloroethane	71-55-6	ND	0.0082	0.022	0.025	< 0.022	
1,1,2-Trichloroethane	79-00-5	ND	0.0070	0.018	0.025	< 0.018	
Trichloroethene	79-01-6	ND	0.0065	0.017	0.025	< 0.017	
Trichlorofluoromethane	75-69-4	ND	0.0068	0.018	0.050	< 0.018	
1,2,3-Trichloropropane	96-18-4	ND	0.010	0.026	0.025	< 0.025	
1,2,4-Trimethylbenzene	95-63-6	ND	0.0029	0.0076	0.025	< 0.0076	

VOST Analytical Results Summary (Continued)
Table A-11. Run 2, VOST Tenax® Tube Pair Field Blank

Analyte	CAS Registry Number	Tenax <sup>®</sup> Tube Pair Field Blank (µg/Sample)						
		Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result <sup>5</sup>	Flag <sup>6</sup>	
1,3,5-Trimethylbenzene	108-67-8	ND	0.0019	0.0050	0.025	< 0.0050		
Vinyl chloride	75-01-4	ND	0.0025	0.0066	0.025	< 0.0066		
m-Xylene & p-Xylene	136777-61-2	ND	0.025	0.066	0.050	< 0.050		
o-Xylene	95-47-6	ND	0.0025	0.0066	0.025	< 0.0066		

### Surrogate Recoveries:

50-150%
50-150%
50-150%
50-150%
50-150%

### BECHTEL BWXT IDAHO, LLC (BBWI)

INTEC HLLWE Effluent Gas Emissions Inventory

Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### VOST Analytical Results Summary (Continued) Table A-11. Run 2, VOST Tenax® Tube Pair Field Blank

#### Sample Collection and Analysis Dates:

	Date		
Date(s) Collected:	June 21, 2001		
Date(s) of Extraction:	July 02, 2001		
Date(s) of Analysis:	July 02, 2001		

#### Sample Collection. Preparation. and Analysis Dates:

SW-846 Method 0031:

"Sampling Method for Volatile Organic Compounds (SMVOC)"

SW-846 Method 5041A:

"Analysis for Desorption of Sorbent Cartridges from Volatile Organic Sampling Train (VOST)"

SW-846 Method 8260B: "Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)"

#### VOST Tentatively Identified Compound (TIC) Summary

TIC <sup>7</sup>	CAS Number	Approximate Retention Time (min.)	Sample Result (μg)	TIC Flag <sup>6</sup>
Hexane, 3-methyl-	589-34-4	4.46	0.028	N,J,M

#### Footnotes:

- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- The RL is the laboratory Reporting Limit (RL).
- Based on the selection rules, the **bolded** value is the value or default value assigned to the analyte using the following guidelines:
  - ♦ When the analytical result is greater than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the result selected by boldface type is the RDL.
  - ♦ When the analytical result is not detected down to the MDL, the result selected by boldface type is the RDL.
  - It should be noted that when the RDL is selected using the guidelines above, but the RL is less than the RDL, the RL is included as the "Risk Result".
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - An "E" flag indicates that the result exceeded the upper calibration range. The analytical result is therefore an estimated value
  - ♦ A "J" flag indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" flag indicates that this compound was found in the associated laboratory method blank. Under these conditions this value is regarded as an estimated value.
  - A "Y" flag indicates that this compound is an indistinguishable isomer as a tentatively identified compound (TIC).
  - An "N" flag indicates that there is presumptive evidence that this compound is present in the sample based on spectral evidence.
  - An "M" flag indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
  - A "D" flag indicates that this result was obtained by a dilution of the sample. The original analysis yielded an analytical result that exceeded the calibration range.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data.

### **VOST Analytical Results Summary** Table A-12. Run 2, VOST Anasorb 747 Tube Field Blank

Field Sample Name: Sample Description: Volatile Organic Sampling Train (VOST)

VOST Anasorb 747 Tube Field Blank for Volatile Organic Compounds Analysis

Field Sample Number(s): STL Sample Number(s).

A-3393 H1F250144-020

	CAS Registry				747 Tube Blank imple)		
Analyte	Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL⁴	Risk Result <sup>5</sup>	Flag <sup>6</sup>
Acetone	67-64-1	0.034	0.024	0.063	0.10	< 0.063	J,B
Acrylonitrile	107-13-1	ND	0.11	0.29	0.50	< 0.29	
Benzene	71-43-2	ND	0.0064	0.017	0.025	< 0.017	
Bromobenzene	108-86-1	ND	0.0039	0.010	0.025	< 0.010	
Bromochloromethane	74-97-5	ND	0.0056	0.015	0.025	< 0.015	
Bromodichloromethane	75-27-4	ND	0.0042	0.011	0.025	< 0.011	
Bromoform	75-25-2	ND	0.0074	0.019	0.025	< 0.019	
Bromomethane	74-83-9	ND	0.0059	0.015	0.050	< 0.015	***************************************
2-Butanone	78-93-3	ND	0.038	0.10	0.10	< 0.10	
n-Butylbenzene	104-51-8	ND	0.0061	0.016	0.025	< 0.016	
sec-Butylbenzene	135-98-8	ND	0.0032	0.0084	0.025	< 0.0084	
tert-Butylbenzene	98-06-6	ND	0.0062	0.016	0.025	< 0.016	
Carbon disulfide	75-15-0	ND	0.0070	0.018	0.025	< 0.018	
Carbon tetrachloride	56-23-5	ND	0.0069	0.018	0.025	< 0.018	
Chlorobenzene	108-90-7	ND	0.0032	0.0084	0.025	< 0.0084	
Chlorodibromomethane	124-48-1	ND	0.0056	0.015	0.025	< 0.015	
Chloroethane	75-00-3	ND	0.0068	0.018	0.050	< 0.018	
Chloroform	67-66-3	ND	0.0070	0.018	0.025	< 0.018	
Chloromethane	74-87-3	ND	0.0048	0.013	0.050	< 0.013	
2-Chlorotoluene	95-49-8	ND	0.0018	0.0047	0.025	< 0.0047	
4-Chlorotoluene	106-43-4	ND	0.0018	0.0047	0.025	< 0.0047	
1,2-Dibromo-3-chloropropane	96-12-8	ND	0.011	0.029	0.050	< 0.029	
1,2-Dibromoethane	106-93-4	ND	0.0075	0.020	0.025	< 0.020	
Dibromomethane	74-95-3	ND	0.0064	0.017	0.025	< 0.017	
1,2-Dichlorobenzene	95-50-1	ND	0.0077	0.020	0.025	< 0.020	
1,3-Dichlorobenzene	541-73-1	ND	0.0038	0.010	0.025	< 0.010	
1,4-Dichlorobenzene	106-46-7	ND	0.0055	0.014	0.025	< 0.014	
Dichlorodifluoromethane	75-71-8	ND	0.0051	0.013	0.025	< 0.013	

## VOST Analytical Results Summary (Continued) Table A-12. Run 2, VOST Anasorb 747 Tube Field Blank

	CAS Registry			Field	747 Tube Blank ample)		
Analyte	Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL⁴	Risk Result <sup>5</sup>	Flag <sup>6</sup>
1,1-Dichloroethane	75-34-3	ND	0.0064	0.017	0.025	< 0.017	
1,2-Dichloroethane	107-06-2	ND	0.0066	0.017	0.025	< 0.017	
1,1-Dichloroethene	75-35-4	ND	0.0067	0.018	0.025	< 0.018	
cis-1,2-Dichloroethene	156-59-2	ND	0.0062	0.016	0.025	< 0.016	
trans-1,2-Dichloroethene	156-60-5	ND	0.0074	0.019	0.025	< 0.019	
1,2-Dichloropropane	78-87-5	ND	0.0049	0.013	0.025	< 0.013	
1,3-Dichloropropane	142-28-9	ND	0.0073	0.019	0.025	< 0.019	
2,2-Dichloropropane	594-20-7	ND	0.0070	0.018	0.025	< 0.018	
1,1-Dichloropropene	563-58-6	ND	0.0077	0.020	0.025	< 0.020	
cis-1,3-Dichloropropene	10061-01-5	ND	0.0046	0.012	0.025	< 0.012	
trans-1,3-Dichloropropene	10061-02-6	ND	0.0059	0.015	0.025	. < 0.015	
Ethylbenzene	100-41-4	ND	0.0035	0.0092	0.025	< 0.0092	
Hexachlorobutadiene	87-68-3	ND	0.012	0.031	0.025	< 0.025	
2-Hexanone	591-78-6	ND	0.024	0.063	0.10	< 0.063	
Isopropylbenzene	98-82-8	ND	0.0024	0.0063	0.025	< 0.0063	
p-Isopropyltoluene	99-87-6	ND	0.0044	0.012	0.025	< 0.012	***************************************
Methylene chloride	75-09-2	0.027	0.016	0.042	0.025	0.027	В
4-Methyl-2-pentanone	108-10-1	ND	0.027	0.071	0.10	< 0.071	
Naphthalene	91-20-3	ND	0.014	0.037	0.025	< 0.025	
n-Propylbenzene	103-65-1	ND	0.0021	0.0055	0.025	< 0.0055	
Styrene	100-42-5	ND	0.0026	0.0068	0.025	< 0.0068	
1,1,2-Tetrachloroethane	630-20-6	ND	0.0037	0.0097	0.025	< 0.0097	
1,1,2,2-Tetrachloroethane	79-34-5	ND	0.0097	0.025	0.025	< 0.025	
Tetrachloroethene	127-18-4	ND	0.0062	0.016	0.025	< 0.016	
Toluene	108-88-3	0.12	0.0025	0.0066	0.025	0.12	
1,2,3-Trichlorobenzene	87-61-6	ND	0.014	0.037	0.025	< 0.025	
1,2,4-Trichlorobenzene	120-82-1	ND	0.013	0.034	0.025	< 0.025	***************************************
1,1,1-Trichloroethane	71-55-6	ND	0.0082	0.022	0.025	< 0.022	
1,1,2-Trichloroethane	79-00-5	ND	0.0070	0.018	0.025	< 0.018	***************************************
Trichloroethene	79-01-6	ND	0.0065	0.017	0.025	< 0.017	
Trichlorofluoromethane	75-69-4	ND	0.0068	0.018	0.050	< 0.018	
1,2,3-Trichloropropane	96-18-4	ND	0.010	0.026	0.025	< 0.025	.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
1,2,4-Trimethylbenzene	95-63-6	ND	0.0029	0.0076	0.025	< 0.0076	

VOST Analytical Results Summary (Continued)
Table A-12. Run 2, VOST Anasorb 747 Tube Field Blank

Analyte	CAS Registry		Anasorb 747 Tube Field Blank (μg/Sample)					
	Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result <sup>5</sup>	Flag <sup>6</sup>	
1,3,5-Trimethylbenzene	108-67-8	ND	0.0019	0.0050	0.025	< 0.0050		
Vinyl chloride	75-01-4	ND	0.0025	0.0066	0.025	< 0.0066		
m-Xylene & p-Xylene	136777-61-2	ND	0.025	0.066	0.050	< 0.050		
o-Xylene	95-47-6	ND	0.0025	0.0066	0.025	< 0.0066		

### Surrogate Recoveries:

,2-Dichloroethane-d <sub>4</sub>	Percent Recovery (%)	Laboratory Recovery Limits (%)	
Dibromofluoromethane	77%	50-150%	
1,2-Dichloroethane-d <sub>4</sub>	68%	50-150%	
Toluene-d <sub>8</sub>	99%	50-150%	
Bromofluorobenzene	73%	50-150%	

INTEC HLLWE Effluent Gas Emissions Inventory

Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

## VOST Analytical Results Summary (Continued) Table A-12. Run 2, VOST Anasorb 747 Tube Field Blank

### Sample Collection and Analysis Dates:

	Date
Date(s) Collected:	June 21, 2001
Date(s) of Extraction:	July 02, 2001
Date(s) of Analysis:	July 02, 2001

### Sample Collection. Preparation. and Analysis Dates:

SW-846 Method 0031:

"Sampling Method for Volatile Organic Compounds (SMVOC)"

SW-846 Method 5041A: SW-846 Method 8260B:

"Analysis for Desorption of Sorbent Cartridges from Volatile Organic Sampling Train (VOST)"

"Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)"

### VOST Tentatively Identified Compound (TIC) Summary

TIC <sup>7</sup>	CAS Number	Approximate Retention Time (min.)	Sample Result (µg)	TIC Flag <sup>6</sup>
Cyclohexane, methyl-	108-87-2	5.32	0.17	N,J,M
Hexane, 2,4-dimethyl-	589-43-5	5.40	0.16	N,J,M
Cyclopentane, ethyl-	1640-89-7	5.49	0.044	N,J,M
Pentane, 2,3-dimethyl-	565-59-3	4.36	0.30	N,J,M
Pentane, 3,3-dimethyl-	562-49-2	4.13	0.064	N,J,M
Hexane, 2-methyl-	591-76-4	4.30	0.26	N,J,M

### Footnotes:

- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- The RL is the laboratory Reporting Limit (RL).
- Based on the selection rules, the **bolded** value is the value or default value assigned to the analyte using the following guidelines:
  - ♦ When the analytical result is greater than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - ♦ When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the result selected by boldface type is the RDL.
  - When the analytical result is not detected down to the MDL, the result selected by boldface type is the RDL.
  - It should be noted that when the RDL is selected using the guidelines above, but the RL is less than the RDL, the RL is included as the "Risk Result".
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - An "E" flag indicates that the result exceeded the upper calibration range. The analytical result is therefore an estimated value.
  - ♦ A "J" flag indicates that this compound was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
  - A "B" flag indicates that this compound was found in the associated laboratory method blank. Under these conditions this value is regarded as an estimated value.
  - A "Y" flag indicates that this compound is an indistinguishable isomer as a tentatively identified compound (TIC).
  - An "N" flag indicates that there is presumptive evidence that this compound is present in the sample based on spectral evidence.
  - An "M" flag indicates that this result was measured against the nearest internal standard and assumed a response factor of one (1).
  - A "D" flag indicates that this result was obtained by a dilution of the sample. The original analysis yielded an analytical result that exceeded the calibration range.
- The tentatively identified compounds (TICs) were identified by conducting a mass spectral library search using the NBS library of data.

### M5 Particulate and Anion Train - Run 1 **Analytical Results Summary** Table A-13. HLLWE Run ID: 0050-STRT-1

Field Sample Name:

M5 HCl/Cl2 and Particulate Train

Sample Description:

Particulate Filter and Acetone Probe Rinse Samples for Particulate Determination

Field Sample Number(s):

A-3308 and A-3309

STL Sample No.:

H1F210104-001 and H1F210104-002

Analyte	Particulate Filter	Acetone Probe Rinse	Particulate <sup>1</sup>
	Particulate Weight	Particulate Weight	Total
	(µg)	(µg)	(Total µg of Particulate)
Particulate	100 B	3,400	3,500 B

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train

Sample Description:

0.1N H<sub>2</sub>SO<sub>4</sub> Impinger Solution for Chloride, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3310

STL Sample No.:

H1F210104-003

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	$RDL^4$	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chloride (as HCl) <sup>8</sup>	1.9	1.7	4.5	2.9	< 2.9	В
Fluoride (as HF) <sup>9</sup>	ND	0.075	0.20	1.5	< 0.20	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	11	0.072	0.19	0.72	11	
Nitrite (as HNO <sub>2</sub> ) <sup>11</sup>	ND	0.073	0.19	0.73	< 0.19	

### M5 Particulate and Anion Train - Run 1 **Analytical Results Summary (Continued)** Table A-13. HLLWE Run ID: 0050-STRT-1

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train

Sample Description:

0.1N NaOH Impinger Solution for Chlorine, Fluoride, Nitrate, and Nitrite Analysis

Flag <sup>7</sup>

Field Sample Number(s):

A-3311 STL Sample No.:

H1F210104-004

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>
Chlorine (as Cl <sub>2</sub> ) 12	ND	0.16	0.42	0.29	< 0.29

	Chlorine (as Cl <sub>2</sub> ) <sup>12</sup>	ND	0.16	0.42	0.29	< 0.29	
	Fluoride (as HF) <sup>9</sup>	ND	0.075	0.20	1.5	< 0.20	
	Nitrate (as HNO <sub>3</sub> ) 10	0.85	0.014	0.037	0.14	0.85	
,	Nitrite (as HNO <sub>2</sub> ) 11	1.4	0.073	0.19	0.73	1.4	

### Sample Collection and Analysis Dates:

	Particulate Acetone Filter Probe Rinse		0.1N H <sub>2</sub> SO <sub>4</sub> Impinger Composite	0.1N NaOH Impinger Composite	
Date(s) Collected:	June 07, 2001	June 07, 2001	June 07, 2001	June 07, 2001	
Date(s) of Preparation- Analysis:	June 22-26, 2001	June 25-26, 2001	June 29, 2001	July 02, 2001	

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056: "Determination of Inorganic Anions by Ion Chromatography"

### BECHTEL BWXT IDAHO, LLC (BBWI) INTEC HLLWE Effluent Gas Emissions Inventory

Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### M5 Particulate and Anion Train - Run 1 Analytical Results Summary (Continued) Table A-13. HLLWE Run ID: 0050-STRT-1

### Footnotes:

1 The Total M-5 Particulate result is the sum of the acetone probe rinse particulate weight and the particulate filter particulate weight.

<sup>2</sup> This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the method detection limit (MDL).

<sup>3</sup> This value is the laboratory MDL derived according to requirements outlined in 40 CFR Part 136, Appendix B. The MDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL). The RDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

<sup>5</sup> The RL is the laboratory Reporting Limit (RL). The RL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The **bolded** value for each analyte is the value or default value assigned to the analyte. This value was determined using the following guidelines:

- When the analytical result is greater than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
- When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
- ♦ When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the default value is the RDL.
- When the analytical result is not detected down to the MDL, the "Risk Result" is the RDL.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is used as the risk result.
- <sup>7</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.
     A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.
- The calculation of the total milligrams (mg) of HCl in the Impinger Composite is as follows:

Total mg (HCI) = mg of Cl<sup>-</sup> 
$$\times \frac{(36.46 \text{ mg HCl})}{(35.45 \text{ mg Cl}^-)}$$

Where: 36.46 = the molecular weight of HCl in mg/mg-mole and 35.45 = the atomic weight of Cl' in mg/mg-mole.

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### M5 Particulate and Anion Train - Run 1 Analytical Results Summary (Continued) Table A-13. HLLWE Run ID: 0050-STRT-1

The calculation of the total milligrams (mg) of HF in Impinger Composite is as follows:

Total mg(HF) = mg of F<sup>-</sup> 
$$\times \frac{(20.01 \text{ mgHF})}{(19.00 \text{ mgF}^-)}$$

Where: 20.01 = the molecular weight of HF in mg/mg-mole and 19.00 = the atomic weight of F in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>3</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>3</sub>) = mg of NO<sub>3</sub><sup>-</sup> × 
$$\frac{(63.01 \text{ mg HNO}_3)}{(62.00 \text{ mg NO}_3^-)}$$

Where: 63.01 = the molecular weight of HNO<sub>3</sub> in mg/mg-mole and 62.00 = the molecular weight of NO<sub>3</sub> in mg/mg-mole.

11 The calculation of the total milligrams (mg) of HNO<sub>2</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>2</sub>) = mg of NO<sub>2</sub><sup>-</sup> × 
$$\frac{(47.01 \text{ mg HNO}_2)}{(46.01 \text{ mg NO}_2^-)}$$

Where: 47.01 = the molecular weight of HNO<sub>2</sub> in mg/mg-mole and 46.01 = the molecular weight of NO<sub>2</sub> in mg/mg-mole.

No additional calculation was required to be performed to obtain the total milligrams (mg) of Cl<sub>2</sub> in the NaOH Impinger Composite. Note that the NaOH Impingers were treated with NaS<sub>2</sub>O<sub>3</sub> prior to the analysis of chlorine.

### M5 Particulate and Anion Train - Run 3 **Analytical Results Summary** Table A-14. HLLWE Run ID: 0050-END-1

Field Sample Name:

M5 HCl/Cl2 and Particulate Train

Sample Description:

Particulate Filter and Acetone Probe Rinse Samples for Particulate Determination

Field Sample Number(s):

A-3338 and A-3339

STL Sample No.:

H1F210104-018 and H1F210104-019

Analyte	Particulate Filter	Acetone Probe Rinse	Particulate <sup>1</sup>
	Particulate Weight	Particulate Weight	Total
	(µg)	(µg)	(Total μg of Particulate)
Particulate	300 B	2,600	2,900 В

Field Sample Name:

M5 HCl/Cl2 and Particulate Train

Sample Description:

0.1N H<sub>2</sub>SO<sub>4</sub> Impinger Solution for Chloride, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3340

STL Sample No.:

H1F210104-020

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chloride (as HC!) <sup>8</sup>	2.1	1.7	4.4	2.9	< 2.9	В
Fluoride (as HF) <sup>9</sup>	ND	0.075	0.20	1.5	< 0.20	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	9.8	0.072	0.19	0.72	9.8	
Nitrite (as HNO <sub>2</sub> ) 11	ND	0.073	0.19	0.73	< 0.19	

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### M5 Particulate and Anion Train - Run 3 **Analytical Results Summary (Continued)** Table A-14. HLLWE Run ID: 0050-END-1

Field Sample Name:

M5 HCl/Cl2 and Particulate Train

Sample Description:

0.1N NaOH Impinger Solution for Chlorine, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3341

H1F210104-021 STL Sample No.:

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chlorine (as Cl <sub>2</sub> ) <sup>12</sup>	0.42	0.32	0.85	0.57	< 0.57	В
Fluoride (as HF) <sup>9</sup>	ND	0.075	0.20	1.5	< 0.20	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	0.61	0.014	0.038	0.14	0.61	
Nitrite (as HNO <sub>2</sub> ) 11	3.6	0.029	0.076	0.29	3.6	

### Sample Collection and Analysis Dates:

	Particulate Filter	Acetone Probe Rinse	0.1N H <sub>2</sub> SO <sub>4</sub> Impinger Composite	0.1N NaOH Impinger Composite	
Date(s) Collected: Date(s) of Preparation-	June 07, 2001  June 22-26, 2001	June 07, 2001 June 25-26, 2001	June 07, 2001  June 29, 2001	June 07, 2001 July 02, 2001	
Analysis:	June 22-20, 2001	Julie 25-20, 2001	June 27, 2001	July 02, 2001	

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056: "Determination of Inorganic Anions by Ion Chromatography"

### M5 Particulate and Anion Train - Run 3 Analytical Results Summary (Continued) Table A-14. HLLWE Run ID: 0050-END-1

### Footnotes:

The Total M-5 Particulate result is the sum of the acetone probe rinse particulate weight and the particulate filter particulate weight.

This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the method detection limit (MDL).

This value is the laboratory MDL derived according to requirements outlined in 40 CFR Part 136, Appendix B. The MDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL). The RDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

<sup>5</sup> The RL is the laboratory Reporting Limit (RL). The RL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The **bolded** value for each analyte is the value or default value assigned to the analyte. This value was determined using the following guidelines:

• When the analytical result is greater than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.

• When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.

• When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the default value is the RDL.

• When the analytical result is not detected down to the MDL, the "Risk Result" is the RDL.

• It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is used as the risk result.

<sup>7</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:

A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.
 A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.

The calculation of the total milligrams (mg) of HCl in the Impinger Composite is as follows:

Total mg (HCl) = mg of Cl<sup>-</sup> 
$$\times \frac{(36.46 \text{ mg HCl})}{(35.45 \text{ mg Cl}^-)}$$

Where: 36.46 = the molecular weight of HCl in mg/mg-mole and 35.45 = the atomic weight of Cl in mg/mg-mole.

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### M5 Particulate and Anion Train - Run 3 Analytical Results Summary (Continued) Table A-14. HLLWE Run ID: 0050-END-1

<sup>9</sup> The calculation of the total milligrams (mg) of HF in Impinger Composite is as follows:

Total mg(HF) = mg of F 
$$\times \frac{(20.01 \text{ mgHF})}{(19.00 \text{mgF})}$$

Where: 20.01 = the molecular weight of HF in mg/mg-mole and 19.00 = the atomic weight of F in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>3</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>3</sub>) = mg of NO<sub>3</sub><sup>-</sup> × 
$$\frac{(63.01 \text{ mg HNO}_3)}{(62.00 \text{ mg NO}_3^-)}$$

Where: 63.01 = the molecular weight of HNO<sub>3</sub> in mg/mg-mole and 62.00 = the molecular weight of NO<sub>3</sub> in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>2</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>2</sub>) = mg of NO<sub>2</sub><sup>-</sup> × 
$$\frac{(47.01 \text{ mg HNO}_2)}{(46.01 \text{ mg NO}_2)}$$

Where: 47.01 = the molecular weight of HNO<sub>2</sub> in mg/mg-mole and 46.01 = the molecular weight of NO<sub>2</sub> in mg/mg-mole.

No additional calculation was required to be performed to obtain the total milligrams (mg) of Cl<sub>2</sub> in the NaOH Impinger Composite. Note that the NaOH Impingers were treated with NaS<sub>2</sub>O<sub>3</sub> prior to the analysis of chlorine.

## M5 Particulate and Anion Train - Run 2 Analytical Results Summary Table A-15. HLLWE Run ID: 0050-STRT-2

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train

Sample Description:

Particulate Filter and Acetone Probe Rinse Samples for Particulate Determination

Field Sample Number(s):

A-3312 and A-3313

STL Sample No.:

H1F210104-005 and H1F210104-006

Analyte	Particulate Filter	Acetone Probe Rinse	Particulate <sup>1</sup>
	Particulate Weight	Particulate Weight	Total
	(μg)	(µg)	(Total µg of Particulate)
Particulate	600	500	1,100

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train

Sample Description:

0.1N H<sub>2</sub>SO<sub>4</sub> Impinger Solution for Chloride, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3314

STL Sample No.:

H1F210104-007

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chloride (as HCl) <sup>8</sup>	1.7	1.6	4.3	2.9	< 2.9	В
Fluoride (as HF) <sup>9</sup>	ND	0.074	0.19	1.5	< 0.19	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	8.4	0.071	0.19	0.71	8.4	
Nitrite (as HNO <sub>2</sub> ) <sup>11</sup>	ND	0.14	0.38	1.4	< 0.38	

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### M5 Particulate and Anion Train - Run 2 **Analytical Results Summary (Continued)** Table A-15. HLLWE Run ID: 0050-STRT-2

Field Sample Name:

M5 HCl/Cl2 and Particulate Train

Sample Description:

0.1N NaOH Impinger Solution for Chlorine, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3315

STL Sample No.:

H1F210104-008

Analyte	Laboratory Result <sup>2</sup> (Total mg)	MDL <sup>3</sup>	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chlorine (as Cl <sub>2</sub> ) <sup>12</sup>	ND	0.16	0.42	0.28	< 0.28	
Fluoride (as HF) <sup>9</sup>	ND	0.074	0.19	1.5	< 0.19	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	0.22	0.014	0.037	0.14	0.22	
Nitrite (as HNO <sub>2</sub> ) 11	1.2	0.029	0.075	0.29	1.2	

### Sample Collection and Analysis Dates:

	Particulate Filter		0.1N H <sub>2</sub> SO <sub>4</sub> Impinger Composite	0.1N NaOH Impinger Composite
Date(s) Collected:	June 11, 2001	June 11, 2001	June 11, 2001	June 11, 2001
Date(s) of Preparation- Analysis:	June 22-26, 2001	June 25-26, 2001	June 29, 2001	July 05, 2001

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056: "Determination of Inorganic Anions by Ion Chromatography"

### M5 Particulate and Anion Train - Run 2 Analytical Results Summary (Continued) Table A-15. HLLWE Run ID: 0050-STRT-2

The Total M-5 Particulate result is the sum of the acetone probe rinse particulate weight and the particulate filter particulate weight.

This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the method detection limit (MDL).

This value is the laboratory MDL derived according to requirements outlined in 40 CFR Part 136, Appendix B. The MDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL). The RDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

<sup>5</sup> The RL is the laboratory Reporting Limit (RL). The RL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The **bolded** value for each analyte is the value or default value assigned to the analyte. This value was determined using the following guidelines:

• When the analytical result is greater than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.

• When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.

• When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the default value is the RDL.

• When the analytical result is not detected down to the MDL, the "Risk Result" is the RDL.

• It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is used as the risk result.

<sup>7</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:

A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.
 A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.

<sup>8</sup> The calculation of the total milligrams (mg) of HCl in the Impinger Composite is as follows:

Total mg (HCI) = mg of Cl<sup>-</sup> 
$$\times \frac{(36.46 \text{ mg HCl})}{(35.45 \text{ mg Cl}^-)}$$

Where: 36.46 = the molecular weight of HCl in mg/mg-mole and 35.45 = the atomic weight of Cl in mg/mg-mole.

### M5 Particulate and Anion Train - Run 2 Analytical Results Summary (Continued) Table A-15. HLLWE Run ID: 0050-STRT-2

The calculation of the total milligrams (mg) of HF in Impinger Composite is as follows:

Total mg(HF) = mg of F<sup>-</sup> 
$$\times \frac{(20.01 \text{ mgHF})}{(19.00 \text{ mgF}^-)}$$

Where: 20.01 = the molecular weight of HF in mg/mg-mole and 19.00 = the atomic weight of F<sup>-</sup> in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>3</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>3</sub>) = mg of NO<sub>3</sub><sup>-</sup> × 
$$\frac{(63.01 \text{ mg HNO}_3)}{(62.00 \text{ mg NO}_3^-)}$$

Where: 63.01 = the molecular weight of HNO<sub>3</sub> in mg/mg-mole and 62.00 = the molecular weight of NO<sub>3</sub> in mg/mg-mole.

11 The calculation of the total milligrams (mg) of HNO<sub>2</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>2</sub>) = mg of NO<sub>2</sub><sup>-</sup> × 
$$\frac{(47.01 \text{ mg HNO}_2)}{(46.01 \text{ mg NO}_2)}$$

Where: 47.01 = the molecular weight of HNO<sub>2</sub> in mg/mg-mole and 46.01 = the molecular weight of NO<sub>2</sub> in mg/mg-mole.

No additional calculation was required to be performed to obtain the total milligrams (mg) of Cl<sub>2</sub> in the NaOH Impinger Composite. Note that the NaOH Impingers were treated with NaS<sub>2</sub>O<sub>3</sub> prior to the analysis of chlorine.

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## M5 Particulate and Anion Train - Run 4 Analytical Results Summary Table A-16. HLLWE Run ID: 0050-END-2

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train

Sample Description:

Particulate Filter and Acetone Probe Rinse Samples for Particulate Determination

Field Sample Number(s):

A-3342 and A-3343

STL Sample No.:

H1F210104-022 and H1F210104-023

Analyte	Particulate Filter	Acetone Probe Rinse	Particulate <sup>1</sup>
	Particulate Weight	Particulate Weight	Total
	(µg)	(µg)	(Total µg of Particulate)
Particulate	700	100 B	800 B

Field Sample Name:

M5 HCl/Cl2 and Particulate Train

Sample Description:

0.1N H<sub>2</sub>SO<sub>4</sub> Impinger Solution for Chloride, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3344

STL Sample No.:

H1F210104-024

Analyte	Laboratory Result <sup>2</sup> (Total mg)	MDL <sup>3</sup>	$\mathrm{RDL}^4$	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chloride (as HCl) <sup>8</sup>	1.8	1.6	4.3	2.9	< 2.9	В
Fluoride (as HF) <sup>9</sup>	ND	0.074	0.19	1.5	< 0.19	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	6.6	0.071	0.19	0.71	6.6	
Nitrite (as HNO <sub>2</sub> ) <sup>11</sup>	ND	0.14	0.38	1.4	< 0.38	

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### M5 Particulate and Anion Train - Run 4 Analytical Results Summary (Continued) Table A-16. HLLWE Run ID: 0050-END-2

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train

Sample Description:

0.1N NaOH Impinger Solution for Chlorine, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s): A-3345

STL Sample No.:

H1F210104-025

Analyte	Laboratory Result <sup>2</sup> (Total mg)	MDL <sup>3</sup>	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chlorine (as Cl <sub>2</sub> ) 12	ND	0.16	0.42	0.29	< 0.29	
Fluoride (as HF) <sup>9</sup>	ND	0.075	0.20	1.5	< 0.20	
Nitrate (as HNO <sub>3</sub> ) 10	0.51	0.014	0.038	0.14	0.51	
Nitrite (as HNO <sub>2</sub> ) 11	2.7	0.015	0.038	0.15	2.7	

### Sample Collection and Analysis Dates:

	Particulate	Acetone	0.1N H <sub>2</sub> SO <sub>4</sub> Impinger	0.1N NaOH Impinger
	Filter	Probe Rinse	Composite	Composite
Date(s) Collected: Date(s) of Preparation- Analysis:	June 11, 2001  June 22-26, 2001	June 11, 2001 June 25-26, 2001	June 11, 2001 June 29, 2001	June 11, 2001 July 05, 2001

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056:

"Determination of Inorganic Anions by Ion Chromatography"

### M5 Particulate and Anion Train - Run 4 Analytical Results Summary (Continued) Table A-16. HLLWE Run ID: 0050-END-2

### Footnotes:

The Total M-5 Particulate result is the sum of the acetone probe rinse particulate weight and the particulate filter particulate weight.

- This value is the laboratory MDL derived according to requirements outlined in 40 CFR Part 136, Appendix B. The MDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.
- <sup>4</sup> The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL). The RDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.
- The RL is the laboratory Reporting Limit (RL). The RL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.
- The **bolded** value for each analyte is the value or default value assigned to the analyte. This value was determined using the following guidelines:
  - ♦ When the analytical result is greater than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
  - ♦ When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the default value is the RDL.
  - When the analytical result is not detected down to the MDL, the "Risk Result" is the RDL.
  - It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is used as the risk result.
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - ♦ A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.

    A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.
- The calculation of the total milligrams (mg) of HCl in the Impinger Composite is as follows:

Total mg (HCI) = mg of CI<sup>-</sup> 
$$\times \frac{(36.46 \text{ mg HCI})}{(35.45 \text{ mg CI}^-)}$$

Where: 36.46 = the molecular weight of HCl in mg/mg-mole and 35.45 = the atomic weight of Cl in mg/mg-mole.

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<sup>&</sup>lt;sup>2</sup> This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the method detection limit (MDL).

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### M5 Particulate and Anion Train - Run 4 Analytical Results Summary (Continued) Table A-16. HLLWE Run ID: 0050-END-2

The calculation of the total milligrams (mg) of HF in Impinger Composite is as follows:

Total mg(HF) = mg of F 
$$\times \frac{(20.01 \,\text{mgHF})}{(19.00 \,\text{mgF})}$$

Where: 20.01 = the molecular weight of HF in mg/mg-mole and

19.00 = the atomic weight of F in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>3</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>3</sub>) = mg of NO<sub>3</sub><sup>-</sup> × 
$$\frac{(63.01 \text{ mg HNO}_3)}{(62.00 \text{ mg NO}_3^-)}$$

Where: 63.01 = the molecular weight of HNO<sub>3</sub> in mg/mg-mole and

62.00 = the molecular weight of NO<sub>3</sub> in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>2</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>2</sub>) = mg of NO<sub>2</sub><sup>-</sup> × 
$$\frac{(47.01 \text{ mg HNO}_2)}{(46.01 \text{ mg NO}_2)}$$

Where: 47.01 = the molecular weight of HNO<sub>2</sub> in mg/mg-mole and

46.01 = the molecular weight of NO<sub>2</sub> in mg/mg-mole.

No additional calculation was required to be performed to obtain the total milligrams (mg) of Cl<sub>2</sub> in the NaOH Impinger Composite. Note that the NaOH Impingers were treated with NaS<sub>2</sub>O<sub>3</sub> prior to the analysis of chlorine.

### Table A-17. M5 Particulate and Anion Train - Run 2 Reagent Blanks **Analytical Results Summary**

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train Reagent Blanks

Sample Description:

Particulate Filter and Acetone Probe Rinse Reagent Blanks for Particulate Determination

Field Sample Number(s):

A-3316 and A-3317

STL Sample No.:

H1F210104-009 and H1F210104-010

Analyte	Particulate Filter	Acetone Probe Rinse	Particulate <sup>1</sup>	
	Particulate Weight	Particulate Weight	Total	
	(µg)	(µg)	(Total µg of Particulate)	
Particulate	500 U	500 U	1,000 U	

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train Reagent Blanks

Sample Description: Field Sample Number(s): 0.1N H<sub>2</sub>SO<sub>4</sub> Impinger Solution Reagent Blanks for Chloride, Fluoride, Nitrate, and Nitrite Analysis

A-3318

STL Sample No.:

H1F210104-011

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chloride (as HCl) <sup>8</sup>	1.6	1.2	3.0	2.0	< 2.0	В
Fluoride (as HF) <sup>9</sup>	ND	0.052	0.14	1.0	< 0.14	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	ND	0.020	0.053	0.20	< 0.053	
Nitrite (as HNO <sub>2</sub> ) <sup>11</sup>	ND	0.10	0.27	1.0	< 0.27	

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## Table A-17. M5 Particulate and Anion Train - Run 2 Reagent Blanks Analytical Results Summary (Continued)

Field Sample Name:

M5  $HCl/Cl_2$  and Particulate Train Reagent Blanks

Sample Description:

0.1N NaOH Impinger Solution Reagent Blanks for Chlorine, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3319

STL Sample No.:

H1F210104-012

Analyte	Laboratory Result <sup>2</sup> (Total mg)	$MDL^3$	RDL⁴	RL <sup>5</sup>	Risk Result (Total mg) <sup>6</sup>	Flag <sup>7</sup>
Chlorine (as Cl <sub>2</sub> ) 12	ND	0.11	0.30	0.20	< 0.20	
Fluoride (as HF) <sup>9</sup>	ND	0.11	0.28	2.1	< 0.28	
Nitrate (as HNO <sub>3</sub> ) <sup>10</sup>	0.012	0.010	0.027	0.10	< 0.027	В
Nitrite (as HNO <sub>2</sub> ) 11	ND	0.010	0.027	0.10	< 0.027	

### Sample Collection and Analysis Dates:

	Particulate	Acetone	0.1N H <sub>2</sub> SO <sub>4</sub> Impinger	0.1N NaOH Impinger
	Filter	Probe Rinse	Composite	Composite
Date(s) Collected: Date(s) of Preparation-Analysis:	June 11, 2001 June 22-26, 2001	June 11, 2001 June 25-26, 2001	June 11, 2001 June 29, 2001	June 11, 2001 July 05, 2001

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056:

"Determination of Inorganic Anions by Ion Chromatography"

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## Table A-17. M5 Particulate and Anion Train - Run 2 Reagent Blanks Analytical Results Summary (Continued)

### Footnotes:

<sup>1</sup> The Total M-5 Particulate result is the sum of the acetone probe rinse particulate weight and the particulate filter particulate weight.

- <sup>2</sup> This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the method detection limit (MDL).
- This value is the laboratory MDL derived according to requirements outlined in 40 CFR Part 136, Appendix B. The MDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL). The RDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.
- <sup>5</sup> The RL is the laboratory Reporting Limit (RL). The RL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.
- The **bolded** value for each analyte is the value or default value assigned to the analyte. This value was determined using the following guidelines:
  - When the analytical result is greater than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the default value is the RDL.
  - When the analytical result is not detected down to the MDL, the "Risk Result" is the RDL.
  - It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is used as the risk result.
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.

    A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.
- The calculation of the total milligrams (mg) of HCl in the Impinger Composite is as follows:

Total mg (HCI) = mg of CI<sup>-</sup> 
$$\times \frac{(36.46 \text{ mg HCI})}{(35.45 \text{ mg CI}^-)}$$

Where: 36.46 = the molecular weight of HCl in mg/mg-mole and 35.45 = the atomic weight of Cl<sup>-</sup> in mg/mg-mole.

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## Table A-17. M5 Particulate and Anion Train - Run 2 Reagent Blanks Analytical Results Summary (Continued)

The calculation of the total milligrams (mg) of HF in Impinger Composite is as follows:

Total mg<sub>(HF)</sub> = mg of F<sup>-</sup> 
$$\times \frac{(20.01 \text{ mgHF})}{(19.00 \text{ mgF}^-)}$$

Where: 20.01 = the molecular weight of HF in mg/mg-mole and

 $19.00 = \text{the atomic weight of } F^{-} \text{ in mg/mg-mole.}$ 

The calculation of the total milligrams (mg) of HNO<sub>3</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>3</sub>) = mg of NO<sub>3</sub><sup>-</sup> × 
$$\frac{(63.01 \text{ mg HNO}_3)}{(62.00 \text{ mg NO}_3^-)}$$

Where: 63.01 = the molecular weight of HNO<sub>3</sub> in mg/mg-mole and

62.00 = the molecular weight of  $NO_3$  in mg/mg-mole.

The calculation of the total milligrams (mg) of HNO<sub>2</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>2</sub>) = mg of NO<sub>2</sub><sup>-</sup> × 
$$\frac{(47.01 \text{ mg HNO}_2)}{(46.01 \text{ mg NO}_2^-)}$$

Where: 47.01 = the molecular weight of HNO<sub>2</sub> in mg/mg-mole and

46.01 = the molecular weight of  $NO_2$  in mg/mg-mole.

No additional calculation was required to be performed to obtain the total milligrams (mg) of Cl<sub>2</sub> in the NaOH Impinger Composite. Note that the NaOH Impingers were treated with NaS<sub>2</sub>O<sub>3</sub> prior to the analysis of chlorine.

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### Table A-18. M5 Particulate and Anion Train Run 2 INTEC Deionized Water Reagent Blank **Analytical Results Summary**

Field Sample Name:

M5 HCl/Cl<sub>2</sub> and Particulate Train INTEC D.I. Water Reagent Blank

Sample Description:

INTEC D.I. Water Reagent Blank for Chloride, Fluoride, Nitrate, and Nitrite Analysis

Field Sample Number(s):

A-3349

STL Sample No.:

H1F210104-013

Analyte	Laboratory Result <sup>1</sup> (Total mg)	MDL <sup>2</sup>	RDL <sup>3</sup>	$\mathbf{RL}^4$	Risk Result (Total mg) <sup>5</sup>	Flag <sup>6</sup>
Chloride (as HCl) <sup>7</sup>	ND	0.060	0.16	0.10	< 0.10	
Fluoride (as HF) <sup>8</sup>	ND	0.0054	0.014	0.11	< 0.014	
Nitrate (as HNO <sub>3</sub> ) <sup>9</sup>	0.0058	0.0052	0.014	0.052	< 0.014	В
Nitrite (as HNO <sub>2</sub> ) <sup>-10</sup>	ND	0.01	0.014	0.052	< 0.014	

### Sample Collection and Analysis Dates:

Date(s) Collected:	June 11, 2001
Date(s) of Preparation-Analysis:	June 29, 2001

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056: "Determination of Inorganic Anions by Ion Chromatography"

### Table A-18. M5 Particulate and Anion Train Run 2 INTEC Deionized Water Reagent Blank Analytical Results Summary (Continued)

### Footnotes:

This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the method detection limit (MDL).

This value is the laboratory MDL derived according to requirements outlined in 40 CFR Part 136, Appendix B. The MDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL). The RDL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The RL is the laboratory Reporting Limit (RL). The RL has been calculated as Cl<sub>2</sub>, HCl, HF, HNO<sub>3</sub>, or HNO<sub>2</sub>, as appropriate.

The **bolded** value for each analyte is the value or default value assigned to the analyte. This value was determined using the following guidelines:

- ♦ When the analytical result is greater than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
- When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the "Risk Result" is the actual analytical result or "hit" determined by the laboratory.
- When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the default value is the RDL.
- When the analytical result is not detected down to the MDL, the "Risk Result" is the RDL.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is used as the risk result.
- <sup>6</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.

    A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.

<sup>7</sup> The calculation of the total milligrams (mg) of HCl in the Impinger Composite is as follows:

Total mg(HCl) = mg of Cl<sup>-</sup> × 
$$\frac{(36.46 \text{ mg HCl})}{(35.45 \text{ mg Cl}^-)}$$

Where: 36.46 = the molecular weight of HCl in mg/mg-mole and 35.45 = the atomic weight of Cl<sup>-</sup> in mg/mg-mole.

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### Table A-18. M5 Particulate and Anion Train Run 2 INTEC Deionized Water Reagent Blank Analytical Results Summary (Continued)

<sup>8</sup> The calculation of the total milligrams (mg) of HF in Impinger Composite is as follows:

Total mg(HF) = mg of F<sup>-</sup> 
$$\times \frac{(20.01 \text{ mgHF})}{(19.00 \text{ mgF}^-)}$$

Where: 20.01 = the molecular weight of HF in mg/mg-mole and

19.00 = the atomic weight of F in mg/mg-mole.

<sup>9</sup> The calculation of the total milligrams (mg) of HNO<sub>3</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>3</sub>) = mg of NO<sub>3</sub><sup>-</sup> × 
$$\frac{(63.01 \text{ mg HNO}_3)}{(62.00 \text{ mg NO}_3^-)}$$

Where: 63.01 = the molecular weight of HNO<sub>3</sub> in mg/mg-mole and

62.00 = the molecular weight of NO<sub>3</sub> in mg/mg-mole.

<sup>10</sup> The calculation of the total milligrams (mg) of HNO<sub>2</sub> in the Impinger Composite is as follows:

Total mg(HNO<sub>2</sub>) = mg of NO<sub>2</sub> × 
$$\frac{(47.01 \text{ mg HNO}_2)}{(46.01 \text{ mg NO}_2)}$$

Where: 47.01 = the molecular weight of HNO<sub>2</sub> in mg/mg-mole and

46.01 = the molecular weight of  $NO_2$  in mg/mg-mole.

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### Table A-19. M5 Particulate and Anion Train Final Acetone Probe Rinse Analytical Results Summary

Field Sample Name: Sample Description:

M5 HCl/Cl<sub>2</sub> and Particulate Train Final Acetone Probe Rinse Final Acetone Probe Rinse for Particulate Determination

Field Sample Number(s):

A-3346

STL Sample No.:

H1G030222-001

Analyte	Final Acetone Probe Rinse Particulate Weight (µg) <sup>I</sup>
Particulate	6,200

Final Acetone Probe Rinse
June 25, 2001
July 03-06, 2001

### Preparation and Analysis Methods:

EPA Method 5:

"Particulate Emissions from Stationary Sources"

SW-846 Method 9056:

"Determination of Inorganic Anions by Ion Chromatography"

### Footnotes:

- 1 This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" flag indicates that this analyte was analyzed for, but was not detected down to the MDL.

A "B" flag indicates that the result for this analyte was below the RL and is therefore considered to be an estimated value.

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### Method 0060 Multi-Metals Train (MMT) Train Total Summary - Run 1 Train Totals **Metallic Analyte Analytical Results Summary** Table A-20. HLLWE Run ID: 0060-STRT-1

Field Sample Name:

Method 0060 Multi-Metals Train (MMT)

Sample Description:

Method 0060 Multi-Metals Train (MMT) Totals for Metals Analysis

Analyte	CAS Registry Number	MMT Front Half Composite <sup>1</sup> (μg)		MMT Back Half Composite <sup>2</sup> (μg)		MMT Sampling Train Totals <sup>3</sup> (Total μg)		Project Specific
		Risk Result	Flag <sup>4</sup>	Risk Result	Flag <sup>4</sup>	Total <sup>5</sup>	Flag	Flag <sup>6</sup>
Aluminum (Al)	7429-90-5	95		42		140		A
Antimony (Sb)	7440-36-0	2.9	В	1.7	В	< 4.6	В	A
Arsenic (As)	7440-38-2	0.92	U	0.66	В	< 1.6	В	Р
Barium (Ba)	7440-39-3	4.8	В	1.9	В	6.7	В	A
Beryllium (Be)	7440-41-7	0.23	В	0.42	U	< 0.65	В	P
Cadmium (Cd)	7440-43-9	0.32	В	0.14	В	< 0.46	В	Α
Chromium (Cr)	7440-47-3	1.7		1.3		3.0		A
Cobalt (Co)	7440-48-4	1.3	В	1.4	В	< 2.7	В	A
Copper (Cu)	7440-50-8	0.94	В	3.4		4.3	В	A
Lead (Pb)	7439-92-1	0.52	U	0.78	В	< 1.3	В	P
Manganese (Mn)	7439-96-5	2.8		16		19		A
Mercury (Hg)	7439-97-6	0.37	В	100		100		A
Nickel (Ni)	7440-02-0	3.8	В	1.2	В	< 5.0	В	A
Selenium (Se)	7782-49-2	2.4		0.78	В	3.2	В	A
Silver (Ag)	7440-22-4	1.9	U	0.71	U	< 2.6		N
Thallium (Tl)	7440-28-0	1.0	U	1.5	U	< 2.5		N
Vanadium (V)	7440-62-2	1.3	U	1.4	U	< 2.7		N
Zinc (Zn)	7440-66-6	18		43		61		A

### Footnotes:

- <sup>1</sup> The MMT Front Half Composite consists of the Quartz Fiber Particulate Filter and the 0.1 N Nitric Acid Probe Rinse Solution.
- The MMT Back Half configuration includes seven (7) impingers. The first impinger remains empty prior to sampling and serves as a moisture knockout impinger. The second and third impingers are charged with 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> and are analyzed with the first impinger contents for all of the metallic analytes including mercury. The fourth impinger is left empty and serves to protect the mercury trapping solution from carryover. Its contents and glassware rinses are analyzed separately for mercury, only. The fifth and sixth impingers contain 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> and are also analyzed for mercury, only. The seventh impinger contains silica gel for a final scrubbing of residual moisture. The MMT back half mercury (Hg) result is the sum of the mercury in the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> fraction, the fourth empty impinger fraction, the 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> fraction, and the 8N HCl impinger rinse fraction. The total micrograms (ug) of mercury in these fractions is included in the MMT back half result.
- <sup>3</sup> The total mass for each metal found in the MMT sampling train consists of the sum of the MMT train's Front Half metals content plus the train's Back Half metals content. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) = Total  $\mu g$  in the Multi-Metals Sampling Train. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g$ 

The MMT Sampling Train Total value was obtained by summing the MMT Front Half results and the MMT Back Half results using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "B" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "B" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MMT Totals are the cumulative set of flags for each train component included as part of the MMT total. A flag attached to a MMT component is carried through to the "MMT Sampling Train Total" column when the associated component analytical result is a significant number in comparison to the MMT Total. That is, if the MMT Total is affected by a MMT component analytical result, the associated flag is carried to the MMT Train Total, but if the MMT Train Total is not affected by a MMT component, the associated flag is not carried through to the MMT Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- <sup>4</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - ♦ A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.

### Method 0060 Multi-Metals Train (MMT) Train Total Summary - Run 3 Train Totals Metallic Analyte Analytical Results Summary Table A-21. HLLWE Run ID: 0060-END-1

Field Sample Name:

Method 0060 Multi-Metals Train (MMT)

Sample Description: Method 0060 Multi-Metals Train (MMT) Totals for Metals Analysis

Number 7429-90-5 7440-36-0	Risk Result	Flag <sup>4</sup>	Risk Result	Flag <sup>4</sup>	5		Project Specific
	230			riag	Total <sup>5</sup>	Flag	Flag <sup>6</sup>
7440-36-0	250		38		270		A
	3.4	В	1.7	В	< 5.1	В	A
7440-38-2	0.92	U	0.66	U	< 1.6		N
7440-39-3	8.5	В	1.6	В	10	В	A
7440-41-7	0.23	U	0.42	В	< 0.65	В	P
7440-43-9	0.68		0.14	В	< 0.82	В	A
7440-47-3	3.4		1.3		4.7		A
7440-48-4	1.3	U	1.4	U	< 2.7		N
7440-50-8	2.2	В	1.4	В	3.6	В	A
7439-92-1	0.52	В	0.78	В	< 1.3	В	A
7439-96-5	5.4		18		23		A
7439-97-6	0.37	В	150		150		A
7440-02-0	4.9		1.1	В	< 6.0	В	A
7782-49-2	1.5		0.63	U	< 2.1		P
7440-22-4	1.9	U	0.71	U	< 2.6		N
7440-28-0	1.0	U	1.4	U	< 2.4		N
7440-62-2	1.3	U	1.4	U	< 2.7		N
7440-66-6	89		30		120		A
	7440-41-7 7440-43-9 7440-47-3 7440-48-4 7440-50-8 7439-92-1 7439-96-5 7439-97-6 7440-02-0 7782-49-2 7440-22-4 7440-28-0 7440-62-2	7440-41-7       0.23         7440-43-9       0.68         7440-47-3       3.4         7440-48-4       1.3         7440-50-8       2.2         7439-92-1       0.52         7439-96-5       5.4         7440-02-0       4.9         7782-49-2       1.5         7440-22-4       1.9         7440-28-0       1.0         7440-62-2       1.3	7440-41-7       0.23       U         7440-43-9       0.68         7440-47-3       3.4         7440-48-4       1.3       U         7440-50-8       2.2       B         7439-92-1       0.52       B         7439-97-6       0.37       B         7440-02-0       4.9         7782-49-2       1.5         7440-22-4       1.9       U         7440-28-0       1.0       U         7440-62-2       1.3       U	7440-41-7       0.23       U       0.42         7440-43-9       0.68       0.14         7440-47-3       3.4       1.3         7440-48-4       1.3       U       1.4         7440-50-8       2.2       B       1.4         7439-92-1       0.52       B       0.78         7439-96-5       5.4       18         7439-97-6       0.37       B       150         7440-02-0       4.9       1.1         7782-49-2       1.5       0.63         7440-22-4       1.9       U       0.71         7440-28-0       1.0       U       1.4         7440-62-2       1.3       U       1.4	7440-41-7         0.23         U         0.42         B           7440-43-9         0.68         0.14         B           7440-47-3         3.4         1.3         U         1.4         U           7440-48-4         1.3         U         1.4         U         U         7440-50-8         2.2         B         1.4         B         B         0.78         B         B         0.78         B         B         150         0.78         B         150         0.37         B         150         0.37         B         150         0.37         B         1.1         B         0.63         U         0.63         U         0.63         U         0.71         U         0.71         U         0.74         U         0.74 <td>7440-41-7       0.23       U       0.42       B       &lt; 0.65</td> 7440-43-9       0.68       0.14       B       < 0.82	7440-41-7       0.23       U       0.42       B       < 0.65	7440-41-7         0.23         U         0.42         B         < 0.65         B           7440-43-9         0.68         0.14         B         < 0.82

#### Footnotes:

- The MMT Front Half Composite consists of the Quartz Fiber Particulate Filter and the 0.1 N Nitric Acid Probe Rinse Solution.
- The MMT Back Half configuration includes seven (7) impingers. The first impinger remains empty prior to sampling and serves as a moisture knockout impinger. The second and third impingers are charged with 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> and are analyzed with the first impinger contents for all of the metallic analytes including mercury. The fourth impinger is left empty and serves to protect the mercury trapping solution from carryover. Its contents and glassware rinses are analyzed separately for mercury, only. The fifth and sixth impingers contain 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> and are also analyzed for mercury, only. The seventh impinger contains silica gel for a final scrubbing of residual moisture. The MMT back half mercury (Hg) result is the sum of the mercury in the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> fraction, the fourth empty impinger fraction, the 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> fraction, and the 8N HCl impinger rinse fraction. The total micrograms (ug) of mercury in these fractions is included in the MMT back half result.
- The total mass for each metal found in the MMT sampling train consists of the sum of the MMT train's Front Half metals content plus the train's Back Half metals content. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) = Total  $\mu g$  in the Multi-Metals Sampling Train. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g$ 

The MMT Sampling Train Total value was obtained by summing the MMT Front Half results and the MMT Back Half results using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "B" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "B" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MMT Totals are the cumulative set of flags for each train component included as part of the MMT total. A flag attached to a MMT component is carried through to the "MMT Sampling Train Total" column when the associated component analytical result is a significant number in comparison to the MMT Total. That is, if the MMT Total is affected by a MMT component analytical result, the associated flag is carried to the MMT Train Total, but if the MMT Train Total is not affected by a MMT component, the associated flag is not carried through to the MMT Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- <sup>4</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.

- ♦ A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- 6 Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.

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### Method 0060 Multi-Metals Train (MMT) Train Total Summary - Run 2 Train Totals Metallic Analyte Analytical Results Summary Table A-22. HLLWE Run ID: 0060-STRT-2

Field Sample Name:

Method 0060 Multi-Metals Train (MMT)

Sample Description:

Method 0060 Multi-Metals Train (MMT) Totals for Metals Analysis

	CAS Registry	MM Front Compo (µg	Half osite <sup>1</sup>	MM Back I Compo (µg	Half site <sup>2</sup> )	MN Samplin Tota (Tota	g Train als³	Project Specific
Analyte	Number	Risk Result	Flag <sup>4</sup>	Risk Result	Flag <sup>4</sup>	Total <sup>5</sup>	Flag	Flag <sup>6</sup>
Aluminum (Al)	7429-90-5	42		34		76		A
Antimony (Sb)	7440-36-0	2.8	В	1.7	В	< 4.5	В	A
Arsenic (As)	7440-38-2	0.92	U	0.66	U	< 1.6		N
Barium (Ba)	7440-39-3	3.5	В	1.6	В	5.1	В	A
Beryllium (Be)	7440-41-7	0.23	В	0.42	U	< 0.65	В	P
Cadmium (Cd)	7440-43-9	0.13	U	0.14	В	< 0.27	В	P
Chromium (Cr)	7440-47-3	1.3		1.2		2.5		A
Cobalt (Co)	7440-48-4	1.3	U	1.4	. U	< 2.7		N
Copper (Cu)	7440-50-8	0.66	В	1.6	В	< 2.3	В	Α
Lead (Pb)	7439-92-1	0.52	U	0.89	В	< 1.4	В	P
Manganese (Mn)	7439-96-5	16		22		38		A
Mercury (Hg)	7439-97-6	0.37	В	110	·	110		A
Nickel (Ni)	7440-02-0	3.3	В	1.2	В	< 4.5	В	Α
Selenium (Se)	7782-49-2	2.0		0.63	В	< 2.6	В	A
Silver (Ag)	7440-22-4	1.9	U	0.71	U	< 2.6		N
Thallium (Tl)	7440-28-0	1.0	U	1.5	U	< 2.5		N
Vanadium (V)	7440-62-2	1.3	U	1.4	U	< 2.7		N
Zinc (Zn)	7440-66-6	10		20		30		A

#### Footnotes:

- The MMT Front Half Composite consists of the Quartz Fiber Particulate Filter and the 0.1 N Nitric Acid Probe Rinse Solution.
- The MMT Back Half configuration includes seven (7) impingers. The first impinger remains empty prior to sampling and serves as a moisture knockout impinger. The second and third impingers are charged with 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> and are analyzed with the first impinger contents for all of the metallic analytes including mercury. The fourth impinger is left empty and serves to protect the mercury trapping solution from carryover. Its contents and glassware rinses are analyzed separately for mercury, only. The fifth and sixth impingers contain 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> and are also analyzed for mercury, only. The seventh impinger contains silica gel for a final scrubbing of residual moisture. The MMT back half mercury (Hg) result is the sum of the mercury in the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> fraction, the fourth empty impinger fraction, the 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> fraction, and the 8N HCl impinger rinse fraction. The total micrograms (ug) of mercury in these fractions is included in the MMT back half result.
- The total mass for each metal found in the MMT sampling train consists of the sum of the MMT train's Front Half metals content plus the train's Back Half metals content. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) = Total  $\mu g$  in the Multi-Metals Sampling Train. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g$ 

The MMT Sampling Train Total value was obtained by summing the MMT Front Half results and the MMT Back Half results using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "B" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "B" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MMT Totals are the cumulative set of flags for each train component included as part of the MMT total. A flag attached to a MMT component is carried through to the "MMT Sampling Train Total" column when the associated component analytical result is a significant number in comparison to the MMT Total. That is, if the MMT Total is affected by a MMT component analytical result, the associated flag is carried to the MMT Train Total, but if the MMT Train Total is not affected by a MMT component, the associated flag is not carried through to the MMT Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- <sup>4</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- 6 Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions.
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions

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### Method 0060 Multi-Metals Train (MMT) Train Total Summary - Run 4 Train Totals Metallic Analyte Analytical Results Summary Table A-23. HLLWE Run ID: 0060-END-2

Field Sample Name:

Method 0060 Multi-Metals Train (MMT)

Sample Description: Method 0060 Multi-Metals Train (MMT) Totals for Metals Analysis

	CAS Registry	Front Compo	MMT Front Half Composite <sup>1</sup> (μg)		MMT Back Half Composite <sup>2</sup> (µg)		MMT Sampling Train Totals <sup>3</sup> (Total µg)	
Analyte	Number	Risk Result	Flag <sup>4</sup>	Risk Result	Flag <sup>4</sup>	Total <sup>5</sup>	Flag	Flag <sup>6</sup>
Aluminum (Al)	7429-90-5	44		29		73		A
Antimony (Sb)	7440-36-0	2.4	В	1.7	В	< 4.1	В	A
Arsenic (As)	7440-38-2	0.92	U	0.66	U	< 1.6		N
Barium (Ba)	7440-39-3	3.6	В	1.1	В	4.7	В	A
Beryllium (Be)	7440-41-7	0.23	U	0.42	В	< 0.65	В	P
Cadmium (Cd)	7440-43-9	0.13	U	0.14	U	< 0.27		N
Chromium (Cr)	7440-47-3	1.2		3.9		5.1		A
Cobalt (Co)	7440-48-4	1.3	U	1.4	U	< 2.7		N
Copper (Cu)	7440-50-8	0.66	U	0.71	В	< 1.4	В	P
Lead (Pb)	7439-92-1	0.52	U	0.68	В	< 1.2	В	P
Manganese (Mn)	7439-96-5	23		49		72		A
Mercury (Hg)	7439-97-6	0.37	В	110		110		A
Nickel (Ni)	7440-02-0	3.0	В	1.1	В	< 4.1	В	A
Selenium (Se)	7782-49-2	1.7		0.63	В	< 2.3	В	A
Silver (Ag)	7440-22-4	1.9	U	0.71	U	< 2.6		N
Thallium (Tl)	7440-28-0	1.0	U	1.4	U	< 2.4		N
Vanadium (V)	7440-62-2	1.3	U	1.4	U	< 2.7		N
Zinc (Zn)	7440-66-6	6.6		10		17		A

#### Footnotes:

- <sup>1</sup> The MMT Front Half Composite consists of the Quartz Fiber Particulate Filter and the 0.1 N Nitric Acid Probe Rinse Solution.
- The MMT Back Half configuration includes seven (7) impingers. The first impinger remains empty prior to sampling and serves as a moisture knockout impinger. The second and third impingers are charged with 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> and are analyzed with the first impinger contents for all of the metallic analytes including mercury. The fourth impinger is left empty and serves to protect the mercury trapping solution from carryover. Its contents and glassware rinses are analyzed separately for mercury, only. The fifth and sixth impingers contain 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> and are also analyzed for mercury, only. The seventh impinger contains silica gel for a final scrubbing of residual moisture. The MMT back half mercury (Hg) result is the sum of the mercury in the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> fraction, the fourth empty impinger fraction, the 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> fraction, and the 8N HCl impinger rinse fraction. The total micrograms (ug) of mercury in these fractions is included in the MMT back half result.
- The total mass for each metal found in the MMT sampling train consists of the sum of the MMT train's Front Half metals content plus the train's Back Half metals content. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) = Total  $\mu g$  in the Multi-Metals Sampling Train. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g$ 

The MMT Sampling Train Total value was obtained by summing the MMT Front Half results and the MMT Back Half results using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- ♦ When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "B" flag is carried through the calculation to the train total.
- ♦ When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "B" flag is carried through the calculation to the train total.
- When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MMT Totals are the cumulative set of flags for each train component included as part of the MMT total. A flag attached to a MMT component is carried through to the "MMT Sampling Train Total" column when the associated component analytical result is a significant number in comparison to the MMT Total. That is, if the MMT Total is affected by a MMT component analytical result, the associated flag is carried to the MMT Train Total, but if the MMT Train Total is not affected by a MMT component, the associated flag is not carried through to the MMT Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- <sup>4</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - ♦ A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.

- A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.
- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.

## Method 0060 Multi-Metals Blank Train (MMT) Train Total Summary Run 2 Blank Train Totals Metallic Analyte Analytical Results Summary Table A-24. HLLWE Run ID: 0060-BT-1

Field Sample Name:

Method 0060 Multi-Metals Blank Train (MMT)

Sample Description: Method 0060 Multi-Metals Blank Train (MMT) Totals for Metals Analysis

9-90-5	(μg) Risk Result	Flag <sup>4</sup>		MMT Back Half Composite <sup>2</sup> (µg)		MMT Sampling Train Totals <sup>3</sup> (Total µg)	
			Risk Result	Flag <sup>4</sup>	Total <sup>5</sup>	Flag	Flag <sup>6</sup>
	46		29		75		A
-36-0	2.5	В	1.7	В	< 4.2	В	A
-38-2	0.92	В	0.66	U	< 1.6	В	P
-39-3	3.4	В	1.3	В	4.7	В	A
-41-7	0.23	U	0.42	U	< 0.65		N
-43-9	0.13	U	0.14	В	< 0.27	В	P
-47-3	0.66	U	0.66	U	< 1.3		N
-48-4	1.3	U	1.4	U	< 2.7		N
-50-8	0.66	U	0.71	В	< 1.4	В	P
-92-1	0.52	U	0.68	В	< 1.2	В	P
-96-5	1.6		3,700		3,700		A
-97-6	0.37	U	2.7	U	< 3.1		N
-02-0	2.8	В	1.2	В	< 4.0	В	A
-49-2	1.7		0.63	U	< 2.3		P
-22-4	1.9	U	0.71	U	< 2.6		N
-28-0	1.0	U	1.5	U	< 2.5		N
-62-2	1.3	U	1.4	U	< 2.7		N
1-66-6	4.7		34		39		A
).	-49-2 -22-4 -28-0 -62-2	-49-2 1.7 -22-4 1.9 -28-0 1.0 -62-2 1.3	-49-2 1.7 -22-4 1.9 U -28-0 1.0 U -62-2 1.3 U	-49-2 1.7 0.63 -22-4 1.9 U 0.71 -28-0 1.0 U 1.5 -62-2 1.3 U 1.4	-49-2 1.7 0.63 U -22-4 1.9 U 0.71 U -28-0 1.0 U 1.5 U -62-2 1.3 U 1.4 U	-49-2	-249-2     1.7     0.63     U     < 2.3

#### Footnotes:

- <sup>1</sup> The MMT Front Half Composite consists of the Quartz Fiber Particulate Filter and the 0.1 N Nitric Acid Probe Rinse Solution.
- The MMT Back Half configuration includes seven (7) impingers. The first impinger remains empty prior to sampling and serves as a moisture knockout impinger. The second and third impingers are charged with 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> and are analyzed with the first impinger contents for all of the metallic analytes including mercury. The fourth impinger is left empty and serves to protect the mercury trapping solution from carryover. Its contents and glassware rinses are analyzed separately for mercury, only. The fifth and sixth impingers contain 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> and are also analyzed for mercury, only. The seventh impinger contains silica gel for a final scrubbing of residual moisture. The MMT back half mercury (Hg) result is the sum of the mercury in the 5% HNO<sub>3</sub>/10% H<sub>2</sub>O<sub>2</sub> fraction, the fourth empty impinger fraction, the 4% KMnO<sub>4</sub>/10% H<sub>2</sub>SO<sub>4</sub> fraction, and the 8N HCl impinger rinse fraction. The total micrograms (ug) of mercury in these fractions is included in the MMT back half result.
- The total mass for each metal found in the MMT sampling train consists of the sum of the MMT train's Front Half metals content plus the train's Back Half metals content. The calculation is as follows:

(Total  $\mu g$  in the Front Half) + (Total  $\mu g$  in the Back Half) = Total  $\mu g$  in the Multi-Metals Sampling Train. Therefore:  $(\mu g) + (\mu g) = \text{Total } \mu g$ 

The MMT Sampling Train Total value was obtained by summing the MMT Front Half results and the MMT Back Half results using the following guidelines:

- When the train component analytical result is greater than the laboratory reporting limit (RL), the result included in the train total is the actual analytical result or "hit" determined by the laboratory.
- When the train component analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result included in the train total is actual analytical result or "hit" determined by the laboratory and the corresponding "B" flag is carried through the calculation to the train total.
- When the train analytical component result is less than the RDL, but greater than the method detection limit (MDL), the result included in the train total is the RDL and the corresponding "B" flag is carried through the calculation to the train total.
- ♦ When the train component analytical result is not detected down to the MDL, the result included in the train total is the RDL and the corresponding "U" flag is carried through the calculation to the train total.
- It should be noted that when the RDL is selected as the default value using the guidelines above, but the RDL is greater than the RL, the RL is included in the train total.

The data flags attached to the MMT Totals are the cumulative set of flags for each train component included as part of the MMT total. A flag attached to a MMT component is carried through to the "MMT Sampling Train Total" column when the associated component analytical result is a significant number in comparison to the MMT Total. That is, if the MMT Total is affected by a MMT component analytical result, the associated flag is carried to the MMT Train Total, but if the MMT Train Total is not affected by a MMT component, the associated flag is not carried through to the MMT Train Total. The combinations of train fractions are conducted following the standard practice of using significant figures found in ASTM E29-93a(1999), "Standard Practice for Using Significant Digits in Test Data to Determine Conformance with Specifications" and Severn Trent Laboratories standard operating procedure number QA-004, "Rounding and Significant Figures".

- <sup>4</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

- When listed, the less than (<) sign indicates that at least one sample fraction result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value for the fraction that is included in the data set total is the default RDL value and the actual value of the total is known to be less than (<) the displayed result.
- Entries in this column are project-specific train total flags that are applied to the run total values and are not standard EPA data flags. These project-specific flags are utilized for the INEEL NWCF HLLWE Effluent Gas Emissions Inventory project and are defined as follows:
  - An "N" flag in this column indicates that the compound was not measured (detected) in any of the sampling train components, or fractions
  - A "P" flag in this column indicates that the compound was measured (detected) in one or more of the train components, or fractions, but not in all of the sampling train fractions.
  - An "A" flag in this column indicates that the compound was measured (detected) in all of the sampling train components, or fractions.

Table A-25. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary Run 2 Front Half Composite Reagent Blank

Field Sample Name:

Method 0060 Multi-Metals Train (MMT) Front Half Composite Reagent Blanks

Sample Description:

Quartz Fiber Particulate Filter and 0.1 N Nitric Acid Probe Rinse Solution Reagent Blanks for Metals (including

Mercury) Analysis

Field Sample ID: STL Sample No.:

A-3297 and A-3298 H1F200234-011

		MMT Front Half Composite Reagent Blank Total μg							
Analyte	CAS Registry Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result for Blank Correction <sup>5</sup>	Flag <sup>6</sup>		
Aluminum (Al)	7429-90-5	43	5.2	14	20	43			
Antimony (Sb)	7440-36-0	2.7	0.60	1.6	6.0	2.7	В		
Arsenic (As)	7440-38-2	0.72	0.35	0.92	1.0	0.72	В		
Barium (Ba)	7440-39-3	3.6	0.35	0.92	20	3.6	В		
Beryllium (Be)	7440-41-7	0.20	0.089	0.23	0.50	0.20	В		
Cadmium (Cd)	7440-43-9	ND	0.050	0.13	0.50	0			
Chromium (Cr)	7440-47-3	0.92	0.25	0.66	1.0	0.92	В		
Cobalt (Co)	7440-48-4	ND	0.50	1.3	5.0	0			
Copper (Cu)	7440-50-8	ND	0.25	0.66	2.5	0			
Lead (Pb)	7439-92-1	ND	0.20	0.52	1.0	0			
Manganese (Mn)	7439-96-5	0.52	0.15	0.39	1.5	0.52	В		
Mercury (Hg)	7439-97-6	ND	0.14	0.37	0.40	0			
Nickel (Ni)	7440-02-0	2.9	0.44	1.2	4.0	2.9	В		
Selenium (Se)	7782-49-2	2.9	0.35	0.92	1.0	2.9			
Silver (Ag)	7440-22-4	ND	0.71	1.9	2.0	0			
Thallium (Tl)	7440-28-0	ND	0.40	1.0	2.0	0			
Vanadium (V)	7440-62-2	ND	0.50	1.3	5.0	0			
Zinc (Zn)	7440-66-6	3.1	0.23	0.60	2.0	3.1			

#### BECHTEL BWXT IDAHO, LLC (BBWI)

INTEC HLLWE Effluent Gas Emissions Inventory

Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### Table A-25. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary (Continued) Run 2 Front Half Composite Reagent Blank

#### Sample Collection and Analysis Dates:

Date(s) Collected:

Date(s) of Digestion (Metals):

Date(s) of Digestion (Mercury):

Date(s) of Analysis (Metals):

Date(s) of Analysis (Mercury):

June 25, 2001

June 28, 2001

Date(s) of Analysis (Mercury):

June 26, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0060: "Determination of Metals in Stack Emissions"

SW-846 Method 7470A: "Mercury in Liquid Waste (Manual Cold Vapor Technique)"
SW-846 Method 6010B: "Inductively Coupled Plasma - Atomic Emission Spectroscopy"

- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- <sup>2</sup> This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- <sup>4</sup> The RL is the laboratory Reporting Limit (RL).
- The column titled "Risk Result for Blank Correction" presents a "0" for elements that were analyzed for but not detected down to the method detection limit (MDL). Method 0060 specifically calls out the criteria that are to be applied to blank correct multi-metals train (MMT) data.
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

### Table A-26. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary Run 2 Back Half Composite Reagent Blank

Field Sample Name:

Method 0060 Multi-Metals Train (MMT) Back Half Impingers Reagent Blank

Sample Description:

5% HNO3 and 10% H2O2 Impingers Reagent Blank for Metals (including Mercury) Analysis

Field Sample ID:

A-3299

STL Sample No.:

H1F200234-012

Analyte		MMT Back Half Composite Reagent Blank Total μg							
	CAS Registry Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result for Blank Correction <sup>5</sup>	Flag <sup>6</sup>		
Aluminum (Al)	7429-90-5	19	3.5	9.2	23	19	В		
Antimony (Sb)	7440-36-0	0.86	0.69	1.8	6.9	0.86	В		
Arsenic (As)	7440-38-2	ND	0.26	0.68	1.1	0			
Barium (Ba)	7440-39-3	0.96	0.34	0.89	23	0.96	В		
Beryllium (Be)	7440-41-7	ND	0.17	0.45	0.57	0			
Cadmium (Cd)	7440-43-9	0.059	0.057	0.15	0.57	0.059	В		
Chromium (Cr)	7440-47-3	0.91	0.26	0.68	1.1	0.91	В		
Cobalt (Co)	7440-48-4	ND	0.57	1.5	5.7	0			
Copper (Cu)	7440-50-8	ND	0.29	0.76	2.9	0			
Lead (Pb)	7439-92-1	0.43	0.27	0.71	1.1	0.43	В		
Manganese (Mn)	7439-96-5	14	0.17	0.45	1.7	14			
Mercury (Hg)	7439-97-6	ND	0.20	0.52	0.80	0			
Nickel (Ni)	7440-02-0	0.56	0.46	1.2	4.6	0.56	В		
Selenium (Se)	7782-49-2	0.28	0.25	0.66	1.1	0.28	В		
Silver (Ag)	7440-22-4	ND	0.29	0.76	2.3	0			
Thallium (Tl)	7440-28-0	ND	0.58	1.5	2.3	0			
Vanadium (V)	7440-62-2	ND	0.57	1.5	5.7	0			
Zinc (Zn)	7440-66-6	4.7	0.51	1.3	2.3	4.7			

#### BECHTEL BWXT IDAHO, LLC (BBWI)

INTEC HLLWE Effluent Gas Emissions Inventory

Idaho National Engineering and Environmental Laboratory (INEEL)

STL Knoxville Project Number: 142503.40

### Table A-26. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary (Continued) Run 2 Back Half Composite Reagent Blank

#### Sample Collection and Analysis Dates:

Date(s) Collected:

Date(s) of Digestion (Metals):

Date(s) of Digestion (Mercury):

Date(s) of Analysis (Metals):

Date(s) of Analysis (Mercury):

June 25, 2001

June 28, 2001

June 26, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0060:

"Determination of Metals in Stack Emissions"

SW-846 Method 7470A:

"Mercury in Liquid Waste (Manual Cold Vapor Technique)"

SW-846 Method 6010B:

"Inductively Coupled Plasma - Atomic Emission Spectroscopy"

- <sup>1</sup> This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- <sup>2</sup> This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- <sup>4</sup> The RL is the laboratory Reporting Limit (RL).
- The column titled "Risk Result for Blank Correction" presents a "0" for elements that were analyzed for but not detected down to the method detection limit (MDL). Method 0060 specifically calls out the criteria that are to be applied to blank correct multi-metals train (MMT) data.
- <sup>6</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

### Table A-27. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary Run 2 Mercury Impinger Composite Reagent Blank

Field Sample Name: Sample Description:

Method 0060 Multi-Metals Train (MMT) Mercury Impingers Reagent Blank 4% KMnO<sub>4</sub> and 10% H<sub>2</sub>SO<sub>4</sub> Impingers Reagent Blank for Mercury (Hg) Analysis

Field Sample ID:

A-3300

STL Sample No.:

H1F200234-013

CAS Registry Analyte Number	MMT Mercury Impinger Composite Total μg						
	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result for Blank Correction <sup>5</sup>	Flag <sup>6</sup>	
Mercury (Hg)	7439-97-6	ND	0.10	0.26	0.42	0	

#### Sample Collection and Analysis Dates:

Date(s) Collected:

June 07, 2001

Date(s) of Digestion:

June 25, 2001

Date(s) of Analysis:

June 26, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0060:

"Determination of Metals in Stack Emissions"

SW-846 Method 7470A:

"Mercury in Liquid Waste (Manual Cold Vapor Technique)"

### Table A-27. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary (Continued) Run 2 Mercury Impinger Composite Reagent Blank

- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- <sup>2</sup> This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- <sup>4</sup> The RL is the laboratory Reporting Limit (RL).
- The column titled "Risk Result for Blank Correction" presents a "0" for elements that were analyzed for but not detected down to the method detection limit (MDL). Method 0060 specifically calls out the criteria that are to be applied to blank correct multi-metals train (MMT) data.
- <sup>6</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

### Table A-28. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary Run 2 HCI Impinger Rinse Solution Reagent Blank

Field Sample Name:

Method 0060 Multi-Metals Train (MMT) Mercury Impinger HCl Rinse Solution Reagent Blank

Sample Description:

8N HCl Impinger Rinse Solution Reagent Blank for Mercury (Hg) Analysis

Field Sample ID:

A-330

STL Sample No.:

H1F200234-014

			8	N HCI Mercury	MT <sup>,</sup> Impinger Rin il μg	se	
Analyte	CAS Registry Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result for Blank Correction <sup>5</sup>	Flag <sup>6</sup>
Mercury (Hg)	7439-97-6	ND	0.23	0.60	0.92	0	

#### Sample Collection and Analysis Dates:

Date(s) Collected:

June 07, 2001

Date(s) of Digestion:

June 25, 2001

Date(s) of Analysis:

June 26, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0060:

"Determination of Metals in Stack Emissions"

SW-846 Method 7470A:

"Mercury in Liquid Waste (Manual Cold Vapor Technique)"

### Table A-28. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary (Continued) Run 2 HCI Impinger Rinse Solution Reagent Blank

- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- <sup>2</sup> This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- <sup>4</sup> The RL is the laboratory Reporting Limit (RL).
- The column titled "Risk Result for Blank Correction" presents a "0" for elements that were analyzed for but not detected down to the method detection limit (MDL). Method 0060 specifically calls out the criteria that are to be applied to blank correct multi-metals train (MMT) data.
- <sup>6</sup> This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an
    estimated value.

### Table A-29. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary Run 2 INTEC Deionized Water Reagent Blank

Field Sample Name:

INTEC Deionized (D.I.) Water Reagent Blank

Sample Description:

INTEC Deionized (D.I.) Water Reagent Blank for Metals (including Mercury) Analysis

Field Sample ID:

A-3348

STL Sample No.:

H1F200234-015

		MMT INTEC Deionized (D.I.) Water Reagent Blank Total µg								
Analyte	CAS Registry Number	Lab Result <sup>1</sup>	MDL <sup>2</sup>	RDL <sup>3</sup>	RL <sup>4</sup>	Risk Result for Blank Correction <sup>5</sup>	Flag <sup>6</sup>			
Aluminum (Al)	7429-90-5	5.2	5.2	14	20	5.2	В			
Antimony (Sb)	7440-36-0	ND	0.60	1.6	6.0	0				
Arsenic (As)	7440-38-2	ND	0.35	0.92	1.0	0				
Barium (Ba)	7440-39-3	ND	0.35	0.92	20	0				
Beryllium (Be)	7440-41-7	0.13	0.089	0.23	0.50	0.13	В			
Cadmium (Cd)	7440-43-9	ND	0.050	0.13	0.50	0				
Chromium (Cr)	7440-47-3	ND	0.25	0.66	1.0	0				
Cobalt (Co)	7440-48-4	. ND	0.50	1.3	5.0	0				
Copper (Cu)	7440-50-8	ND	0.25	0.66	2.5	0				
Lead (Pb)	7439-92-1	ND	0.20	0.52	1.0	0				
Manganese (Mn)	7439-96-5	ND	0.15	0.39	1.5	0				
Mercury (Hg)	7439-97-6	ND	0.14	0.37	0.40	0				
Nickel (Ni)	7440-02-0	ND	0.44	1.2	4.0	0				
Selenium (Se)	7782-49-2	ND	0.35	0.92	1.0	0				
Silver (Ag)	7440-22-4	ND	0.71	1.9	2.0	0				
Thallium (Tl)	7440-28-0	ND	0.40	1.0	2.0	0				
Vanadium (V)	7440-62-2	ND	0.50	1.3	5.0	0				
Zinc (Zn)	7440-66-6	0.93	0.23	0.60	2.0	0.93	В			

### Table A-29. Method 0060 Multi-Metals Train (MMT) Analytical Results Summary (Continued) Run 2 INTEC Deionized Water Reagent Blank

#### Sample Collection and Analysis Dates:

Date(s) Collected:

Date(s) of Digestion (Metals):

Date(s) of Digestion (Mercury):

Date(s) of Analysis (Metals):

Date(s) of Analysis (Mercury):

June 25, 2001

June 28, 2001

June 26, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0060: "Determination of Metals in Stack Emissions"

SW-846 Method 7470A: "Mercury in Liquid Waste (Manual Cold Vapor Technique)" SW-846 Method 6010B: "Inductively Coupled Plasma - Atomic Emission Spectroscopy"

- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- <sup>2</sup> This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- The RL is the laboratory Reporting Limit (RL).
- The column titled "Risk Result for Blank Correction" presents a "0" for elements that were analyzed for but not detected down to the method detection limit (MDL). Method 0060 specifically calls out the criteria that are to be applied to blank correct multi-metals train (MMT) data.
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - ♦ A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

Table A-30. Final (Post-Test) Acetone Probe Rinse and Nitric Acid Probe Rinse Composite **Metallic Analyte Analytical Results Summary** 

Field Sample Name:

Final (Post-Test) Acetone Probe Rinse and Nitric Acid Probe Rinse Composite

Sample Description:

Final (Post-Test) Acetone Probe Rinse and Nitric Acid Probe Rinse Composite Sample for Metals (including

Mercury) Analysis

Field Sample ID:

A-3346 and A-3347 H1G030222-002

STL Sample No.:

	CAS Registry										
Analyte	Number	Lab Result <sup>2</sup>	MDL <sup>3</sup>	RDL <sup>4</sup>	RL <sup>5</sup>	Risk Result <sup>6,7</sup>	Flag <sup>8</sup>				
	<b>540</b> 0 00 5	100	<b></b>	1.4	20.0	120					
Aluminum (Al)	7429-90-5	120	5.2	14			ъ				
Antimony (Sb)	7440-36-0	1.3	0.60	1.6	6.0	< 1.6	В				
Arsenic (As)	7440-38-2	ND	0.35	0.92	1.0	< 0.92					
Barium (Ba)	7440-39-3	3.2	0.35	0.92	20.0	3.2	В				
Beryllium (Be)	7440-41-7	ND	0.089	0.23	0.50	< 0.23					
Cadmium (Cd)	7440-43-9	4.1	0.050	0.13	0.50	4.1					
Chromium (Cr)	7440-47-3	1.7	0.25	0.66	1.0	1.7					
Cobalt (Co)	7440-48-4	0.85	0.50	1.3	5.0	< 1.3	В				
Copper (Cu)	7440-50-8	2.5	0.25	0.66	2.5	2.5					
Lead (Pb)	7439-92-1	3.2	0.20	0.52	1.0	3.2					
Manganese (Mn)	7439-96-5	6.1	0.15	0.39	1.5	6.1					
Mercury (Hg)	7439-97-6	0.77	0.14	0.37	0.40	0.77					
Nickel (Ni)	7440-02-0	3.2	0.44	1.2	4.0	3.2	В				
Selenium (Se)	7782-49-2	ND	0.35	0.92	1.0	< 0.92					
Silver (Ag)	7440-22-4	1.8	0.71	1.9	2.0	< 1.9	В				
Thallium (Tl)	7440-28-0	ND	0.40	1.0	2.0	< 1.0					
Vanadium (V)	7440-62-2	ND	0.50	1.3	5.0	< 1.3					
Zinc (Zn)	7440-66-6	50	0.23	0.60	2.0	50					

### Table A-30. Final (Post-Test) Acetone Probe Rinse and Nitric Acid Probe Rinse Composite Metallic Analyte Analytical Results Summary (Continued)

#### Sample Collection and Analysis Dates:

Date(s) Collected:	June 25, 2001
Date(s) of Digestion (Metals):	July 08, 2001
Date(s) of Digestion (Mercury):	July 08, 2001
Date(s) of Analysis (Metals):	July 28, 2001
Date(s) of Analysis (Mercury):	July 08, 2001

#### Preparation and Analysis Methods:

SW-846 Method 0060: "Determination of Metals in Stack Emissions"

SW-846 Method 7470A: "Mercury in Liquid Waste (Manual Cold Vapor Technique)" SW-846 Method 6010B: "Inductively Coupled Plasma - Atomic Emission Spectroscopy"

### Table A-30. Final (Post-Test) Acetone Probe Rinse and Nitric Acid Probe Rinse Composite Metallic Analyte Analytical Results Summary (Continued)

- Based on the selection rules, the **bolded** value is the value or default value assigned to the analyte and is carried through to the sampling train total, if it is determined to be significant.
- This value is the laboratory sample result. When the analytical result is "ND" or not detected, the laboratory analysis did not detect the analyte down to the MDL.
- This value is the laboratory Method Detection Limit (MDL) derived according to requirements outlined in 40 CFR Part 136, Appendix B.
- <sup>4</sup> The RDL is the Reliable Detection Limit. The RDL is the detection level recommended by EPA's National Research Laboratory in Cincinnati, Ohio, Environmental Monitoring Systems Laboratory (EMSL) in Cincinnati, Ohio, American Chemical Society (ACS) Committee on Environmental Improvement and the Drinking Water Standards Division (DWSD). It is defined as 2.623 times the MDL (2.623 X MDL).
- <sup>5</sup> The RL is the laboratory Reporting Limit (RL).
- Based on the selection rules, the **bolded** value is the value or default value assigned to the analyte using the following guidelines:
  - When the analytical result is greater than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is greater than the reliable detection level (RDL), but less than the laboratory reporting limit (RL), the result selected by boldface type is the actual analytical result or "hit" determined by the laboratory.
  - When the analytical result is less than the RDL, but greater than the method detection limit (MDL), the result selected by boldface type is the RDL.
  - When the analytical result is not detected down to the MDL, the result selected by boldface type is the RDL.
  - It should be noted that when the RDL is selected using the guidelines above, but the RL is less than the RDL, the RL is included as the "Risk Result".
- When listed, the less than (<) sign indicates that the sample result is either a "non-detect" value down to the MDL of the measurement that carries, or an estimated "hit" value that is below the RDL. In either case, the final value is the default RDL value and the actual value is known to be less than (<) the displayed result.
- This flag is the laboratory data flag that corresponds to EPA guidelines. The data flags for these samples are as follows:
  - A "U" qualifier indicates that this analyte was analyzed for, but was not detected down to the MDL.
  - ♦ A "B" flag indicates that this analyte was detected, but at a concentration below the laboratory RL. The analytical result is therefore an estimated value.

# APPENDIX B OFFGAS SAMPLING DATA

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Table B-1. SVOC-STRT-1.

#### **SVOC SAMPLING DATA SHEET FOR HLLWE TESTS**

Site:	LILL VA/E	Offgas Tie-in		Sampling	Location	NAA	N OFC 72	Nozzle No				2.04	Est. DP:	0.45	E. T. L. 185 400
Project:		062-01-0866		Duct ID. ii		IVIA	12	Nozzle No					Est. K:	-	Est. Tstack, °F: 133
Date:	01-1	6/18/2001			essure, in. WG:		-17.5	Pitot No.:	e, III				Est. DH:		Est. vs, ft/s: 25.8  Operator(s): FE, RW, JA
Run No.:		010-STRT-1		Est. O2, 9	<del></del>		20.6	Pitot Coeff	F -		·		Est. DGM		
Run Type		TEST	,	Est CO2,		· · ·	0	Meter Box					Meter Box		
Pbar., in.		25.238		Est. Moist			1.3%	+	110.	-			Pretest		cfm @ 15 in. Hg
Tambient,		60		Impinger			9	Y-factor:		·		1.0328	1 101001	0.004	Pitot: pass
DGM vol.	Goal (m³):	3.00			Goal (ft <sup>3</sup> ):		127.080	Min. endin	g DGM vo	. (ft <sup>3</sup> ):			Post-test	0.002	cfm @ 8 in. Hg
Sampling	Clock	Velocity		Meter	Meter			TEN	MPERATURE	(°F)			Pump		
Time	Time	ΔΡ		ΔΗ	Volume	Heated		Me	eter	<u>, , , , , , , , , , , , , , , , , , , </u>	Impinger	Aux.	Vacuum	%I <sub>i</sub>	COMMENTS
(min.)	(24hr)	(in. WG)		(in. WG)	(cubic feet)	Line	Stack	In	Out	Filter	Exit	(XAD)	(in. Hg)		
0	8:30	0.15	0.387	1.30	230.461	260	133	72	64	263	53	45	8.0		
10	8:40	0.15	0.387	1.30	237.360	261	133	76	65	262	49	44	8.0	102	O2 analyzer =20.5%
20	8:50	0.15	0.387	1.30	244.556	247	133	79	67	264	50	44	8.0	106	O2 analyzer =20.5%
30	9:00	0.15	0.387	1.30	251.562	261	132	82	68	263	46	45	8.0	103	O2 analyzer =20.5%
40	9:10	0.15	0.387	1.30	258.689	261	132	83	70	262	45	45	8.0	105	O2 analyzer =20.5%
50	9:20	0.15	0.387	1.30	265.823	261	132	84	71	261	45	47	8.0	104	O2 analyzer =20.5%
60	9:30	0.15	0.387	1.30	272.975	261	132	84	72	262	46	48	8.0	105	O2 analyzer =20.5%
70	9:40	0.15	0.387	1.30	280.105	261	132	85	75	261	46	49	8.0	104	O2 analyzer =20.5%
80	9:50	0.15	0.387	1.30	287.461	261	132	85	73	264	46	50	8.0	107	O2 analyzer =20.5%
90	10:00	0.15	0.387	1.30	294.590	261	132	86	73	262	47	50	8.0	104	O2 analyzer =20.5%
100	10:10	0.15	0.387	1.30	301.627	261	132	87	74	263	47	50	8.0	102	O2 analyzer =20.6%
110	10:20	0.15	0.387	1.30	308.833	261	132	87	74	262	48	51	8.0	105	O2 analyzer =20.5%
120	10:30	0.15	0.387	1.30	315.990	261	132	88	75	263	48	50	8.0	104	O2 analyzer =20.6%
130	10:40	0.15	0.387	1.30	323.180	261	132	88	75	262	48	51	8.0	104	O2 analyzer =20.6%
140	10:50	0.15	0.387	1.30	330.485	261	132	89	76	261	49	55	8.0	106	O2 analyzer =20.6%
150	11:00	0.15	0.387	1.30	337.600	261	132	89	77	262	49	50	8.0	103	O2 analyzer =20.6%
160	11:10	0.15	0.387	1.30	344.840	261	132	89	77	261	48	49	8.0	105	O2 analyzer =20.5%
170	11:20	0.15	0.387	1.30	352.138	261	132	90	78	262	48	49	8.0	106	O2 analyzer =20.5%
180	11:30	0.15	0.387	1.30	359.336	261	132	90	78	262	49	49	8.0	104	END OF TEST
Total	Total	ΔPavg		Average	Total			Average	Temperat	ures (°F)			Max.	Ave. %I	
180	3:00	0.150	0.387	1.30	128.875	260	132	85	73	262	48	48	8.0	104	

Table B-1. SVOC-STRT-1.

### 0010 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site:		fgas Tie-in	Impinger Box no.: XAD trap Quanterra No.:		9 <b>A-3355</b>				
Run No.:	0010-STRT-1							7	
Component:	XAD	KO-1	Imp-1	Imp-2		Acid Scrub Section	1		
Туре:	trap	short stem	modified	G-S	short stem	modified	modified		
Reagent:	XAD-2	None	Organics	free water	None	2N NaOH	Silica Gel		
Nominal Contents:	20 - 40g, dry	Empty	100 mL	100 mL	Empty	100 mL	300-400g		
Post-test Wt., g:	272.8	544.6	676.9	690.5	573.6	730.8	804.2	Impinger Wt. Gain	
Pre-test Wt., g:	272.1	541.7	677.7	690.8	572.8	732.2	782.0	Impringer vvi. Gain	
Wt. Gain, g:	0.7	2.9	-0.8	-0.3	0.8	-1.4	22.2	24.1	
Post-test Volume:		0.0	100.0	100.0	0.0			Impinger Vol. Gain	
Pre-test Volume:		0.0	100.0	100.0	0.0			impinger voi. Gain	
Volume Gain:		0.0	0.0	0.0	0.0			0.0	
Post-test pH:			6.0	6.0		14.0			

	Filter Lot #	STL-A4023	OF water Lot #	QCLAB-1	NaOH Lot#	000381
O2%_	20.6	_				
CO2%	0.0	_				

Record impinger change-out and other important information below:

Table B-1. SVOC-STRT-1.

Project: 01-1062-01-0866  Run Date: 6/18/2001								
Run Date: Run Identification:	0010-S							
PARAMETER	SYMBOL	UNITS						
Absolute Pressure in the Duct	Pabs	in. Hg	23.951					
Average Duct Gas Temperature	Ts	R	592					
Average Meter Temperature	Tm	R	538					
Average Gas Oxygen Content	Co2,m	%	20.6					
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0					
Total Impinger Weight Gain (water)	Ww	grams	24.1					
Nozzle Area	An	ft²	0.000538					
Duct Area	As	ft²	0.7854					
Sample Volume	VmStd	dscf	110.435					
Sample Volume (SI)	VmStdm	dscm	3.127					
Average Sampling Rate	Qm	dscf/m	0.614					
Volume of Water Vapor	VwStd	scf	1.136					
Volume of Water Vapor (SI)	VwStdm	scm	0.0322					
Moisture Fraction	Bws	-	0.010					
Dry Gas Molecular Weight	Md	g/g-mol	28.82					
Wet Gas Molecular Weight	Ms	g/g-mol	28.71					
Gas Velocity at Nozzle	vn	ft/s	25.8					
Gas Velocity at Nozzle (SI)	vnm	m/s	7.86					
Average Gas Velocity	vncor	ft/s	21.77					
Dry Offgas Flow Rate	Qsd	dscf/h	43,494					
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,231.6					
Actual Offgas Flow Rate	Q	acf/h	61,566					
Intermediate Isokinetic Rate	li	%	104.6					
Final Isokinetic Rate	I	%	104.3					

Table B-1. SVOC-STRT-1.

### **RESULTS**

			ON	CENTRATION				٨	//AS	S FLOW RATE	S	
		Actual		Standard	1	Dry Standard			Г			
		(µg/acm)		(µg/scm)	i	(µg/dscm)		μg/min		grams/sec		lb/h
Acenaphthene	<	2.214e0		3.1e0		3.1e0		6.4e1		1.1e-6		8.5e-
Acenaphthylene	<	2.1e0		3.0e0		3.0e0		6.2e1		1.0e-6		8.2e-
Acetophenone	<,J	1.1e1	<,J	1.6e1	<,J		1 '	3.2e2				4.3e-
Aniline	< .	2.5e1	<	3.5e1	< .	3.5e1	<	7.2e2	<	1.2e-5		9.6e-
Anthracene		2.1e0	<	3.0e0	<	3.0e0		6.2e1		1.0e-6		8.2e-
Benzidine	<	1.5e2	<	2.1e2	<	2.1e2		4.4e3		7.3e-5		5.8e-
Benzoic acid	E	1.1e3		1.6e3		1.6e3		3.2e4		5.4e-4		4.3e-
Benzo(a)anthracene	<	2.7e0	<	3.8e0	<	3.8e0		7.9e1	<_	1.3e-6	•	1.0e-
Benzo(a)pyrene	<	2.9e1	<	4.1e1	<	4.2e1	<	8.5e2		1.4e-5		1.1e-
Benzo(b)fluoranthene	<	6.8e1	<	9.5e1	<	9.6e1	<	2.0e3		3.3e-5	1	2.6e~
Benzo(g,h,i)perylene	<	3.8e1	<	5.4e1	<	5.4e1	<	1.1e3		1.9e-5		1.5e-
Benzo(k)fluoranthene	<	9.7e1	<	1.4e2	<_	1.4e2		2.8e3		4.7e-5		3.7e-
Benzyl alcohol	<	1.3e2	<	1.8e2	<	1.8e2		3.7e3		6.2e-5		4.9e-
bis(2-Chloroethoxy)methane	< <	2.3e0	< <	3.2e0	<	3.2e0 3.5e0		6.6e1 7.2e1		1.1e-6 1.2e-6		8.7e-4 9.6e-4
bis(2-Chloroethyl)ether		2.5e0		3.5e0	1			7.2e1 7.9e2	1			1.0e-
bis(2-Ethylhexyl)phthalate	<,J <	2.7e1	<,J <	3.8e1	<,J			6.2e1	<,J	1.0e-6		8.2e-6
4-Bromophenyl-phenylether	<	2.1e0 2.9e0	<	3.0e0 4.1e0	<	3.0e0 4.2e0		6.2e1 8.5e1	<	1.0e-6 1.4e-6	1	8.2e-4
Butylbenzylphthalate Carbazole	<	2.9e0 2.9e0	<	4.1e0 4.1e0	<	4.2e0 4.2e0		8.5e1		1.4e-6 1.4e-6		1.1e-
Garbazole 4-Chloro-3-methylphenol		2.9e0 3.8e0	<i>'</i>	5.4e0	<	5.4e0	`	1.1e2		1.4e-6		1.5e-
4-Chloroaniline	<	2.0e1	/ · ·	2.8e1	<	2.9e1	~	5.9e2		9.8e-6		7.8e-
2-Chloronaphthalene	<	2.1e0	` <b>'</b>	2.9e0	<	3.0e0		6.1e1		1.0e-6		8.1e-6
2-Chlorophenol	<	2.1e0 2.5e0	<	3.5e0	<	3.5e0	I	7.2e1	1	1.2e-6		9.6e-6
4-Chlorophenyl phenyl ether	<	2.5e0	` <	3.5e0	<	3.5e0		7.2e1		1.2e-6	1	9.6e-6
Chrysene	<	2.7e0	<	3.8e0	<	3.8e0		7.9e1		1.3e-6	-	1.0e-
Di-n-butylphthalate	<,j	2.3e1	<,J	3.2e1	<.J			6.6e2				8.7e-
Di-n-octylphthalate	<,J	3.6e1		5.1e1	<,J			1.1e3	1 '			1.4e-4
Dibenz(a,h)anthracene	<"	3.8e1	<	5.4e1	<"	5.4e1	<"	1.1e3		1.9e-5		1.5e-4
Dibenzofuran	<	2.5e0	<	3.5e0	<	3.5e0	<	7.2e1		1.2e-6		9.6e-6
1,2-Dichlorobenzene	<	2.5e0	<	3.5e0	<	3.5e0	<	7.2e1	<	1.2e-6	<	9.6e-6
1,3-Dichlorobenzene	<	2.7e0	<	3.8e0	<	3.8e0	<	7.9e1	<	1.3e-6	<	1.0e-
1,4-Dichlorobenzene	<,J	3.6e0	<,J	5.1e0	<,J	5.1e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-
3,3'-Dichlorobenzidine	<	2.5e1	<	3.5e1	<	3.5e1	<	7.2e2	<	1.2e-5	<	9.6e-
2,4-Dichlorophenol	<	2.9e0	<	4.1e0	<	4.2e0	<	8.5e1	<	1.4e-6	<	1.1e-
Diethylphthalate	<,J	3.6e0	<,J	5.1e0	<,J	5.1e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-
Dimethyl phthalate	<	2.2e0	<	3.0e0	<	3.1e0	<	6.3e1	<	1.1e-6	<	8.3e-6
2,4-Dimethylphenol	<	1.3e1	<	1.9e1	<	1.9e1		3.9e2		6.5e-6		5.1e-
4,6-Dinitro-2-methylphenol	<	2.9e1	<	4.1e1	<	4.2e1	<	8.5e2	<	1.4e-5	1	1.1e-
2,4-Dinitrophenol	<	6.1e1		8.5e1	<	8.6e1		1.8e3		3.0e-5	1	2.3e-4
2,4-Dinitrotoluene	<	2.9e0	<	4.1e0	<_	4.2e0		8.5e1	<	1.4e-6		1.1e-
2,6-Dinitrotoluene	<	2.7e0	<	3.8e0	<	3.8e0	1	7.9e1	<	1.3e-6	1	1.0e-
1,2-Diphenylhydrazine	<	2.2e0	<	3.1e0	<	3.1e0		6.4e1	<	1.1e-6	1	8.5e-6
Fluoranthene	<	2.3e0	<	3.2e0	<	3.2e0		6.6e1		1.1e-6		8.7e-6
Fluorene	<	2.3e0	<	3.2e0	<	3.2e0	<	6.6e1	<	1.1e-6		8.7e-6
Hexachlorocyclopentadiene	<	3.6e1	<	5.1e1	<	5.1e1	<	1.1e3		1.8e-5		1.4e-4
Hexachlorobenzene	<	2.5e0		3.5e0		3.5e0	<	7.2e1		1.2e-6		9.6e-6
Hexachlorobutadiene	<	3.4e0	<	4.7e0	<	4.8e0	<	9.8e1	,	1.6e-6		1.3e-
Hexachloroethane	<	3.6e0	<	5.1e0	<	5.1e0	<	1.1e2		1.8e-6		1.4e-
Indeno(1,2,3-cd)pyrene	<	3.4e1	<	4.7e1	<	4.8e1	<	9.8e2		1.6e-5		1.3e-4
Isophorone	<	2.3e0	<	3.2e0	<	3.2e0	<	6.6e1		1.1e-6		8.7e-6
2-Methylnaphthalene	<	2.3e0	<	3.2e0	<	3.2e0	<	6.6e1	<	1.1e-6		8.7e-0
2-Methylphenol	<	1.1e1	<	1.5e1	<	1.5e1	<	3.1e2		5.1e-6		4.1e-
3-Methylphenol & 4-Methylphenol	<	7.7e0	<	1.1e1	<	1.1e1	1	2.2e2		3.7e-6		
N-Nitroso-di-n-propylamine	< <	2.5e0	<	3.5e0	<	3.5e0	<	7.2e1	<	1.2e-6		9.6e-
N-Nitrosodimethylamine	1	2.5e0	<	3.5e0	1 -	3.5e0		7.2e1	1	1.2e-6		9.6e-
N-Nitrosodiphenylamine	<	3.4e0	<	4.7e0	<	4.8e0	<	9.8e1	<	1.6e-6		1.3e-
Naphthalene	<	2.5e0	< <	3.5e0 3.5e0	<	3.5e0 3.5e0	<	7.2e1 7.2e1	<	1.2e-6 1.2e-6		9.6e-6 9.6e-6
2-Nitroaniline	<	2.5e0			<				1			
3-Nitroaniline	<	9.3e0	<	1.3e1	,	1.3e1	1	2.7e2		4.5e-6		3.6e-
4-Nitroaniline	<	8.1e0	<	1.1e1	<_	1.2e1	<u></u>	2.4e2	<u>'</u>	3.9e-6	L	3.1e-

Table B-1. SVOC-STRT-1.

Project: 01-1062-01-0866
Run Date: 6/18/2001
Run Identification: 0010-STRT-1 Run loentinication: Run Type: Lab Report Date: Lab Report Status: (preliminary or final) TEST 8/28/2001 Final

#### **RESULTS**

							ASS F	LOW RATE	ES			
	Ad	tual	Sta	ndard		tandard	l					
	(µg	/acm)	(µg	/scm)	(μg/	dscm)	μί	g/min	gra	ms/sec		lb/h
Nitrobenzene	<,J	3.6e0	<,J	5.1e0		5.1e0		1.1e2		1.8e-6		1.4e-5
2-Nitrophenol	<	2.1e1	<	2.9e1		2.9e1		6.0e2		1.0e-5		7.9e-5
4-Nitrophenol	ئ,>			2.4e1	<,J	2.4e1		5.0e2		8.3e-6		6.6e-5
2,2'-Oxybis(1-chloropropane)	. <	3.2e0		4.4e0		4.5e0		9.2e1		1.5e-6		1.2e-5
Pentachlorobenzene	<	2.3e0		3.2e0		3.2e0		6.6e1	<	1.1e-6		8.7e-6
Pentachloronitrobenzene	<	2.5e0	l .	3.5e0	1	3.5e0		7.2e1		1.2e-6		9.6e-6
Pentachlorophenol	<	6.8e1		9.5e1		9.6e1		2.0e3	i	3.3e-5		2.6e-4
Phenanthrene	<	2.2e0		3.1e0		3.1e0		6.4e1		1.1e-6		8.5e-6
Phenol	<	1.7e1	<	2.4e1		2.5e1		5.1e2		8.4e-6		6.7e-5
Pyrene	<	2.3e0	<	3.2e0	<	3.2e0	<	6.6e1		1.1e-6		8.7e-6
Pyridine	<	3.8e0	l .	5.4e0	1	5.4e0		1.1e2		1.9e-6		1.5e-5
1,2,4,5-Tetrachlorobenzene	<	2.5e0	<	3.5e0		3.5e0		7.2e1		1.2e-6		9.6e-6
1,2,4-Trichlorobenzene	<	2.7e0	<	3.8e0	<	3.8e0		7.9e1		1.3e-6		1.0e-5
2,4,5-Trichlorophenol	<	5.6e0	l .	7.9e0		8.0e0		1.6e2		2.7e-6		2.2e-5
2,4,6-Trichlorophenol <b>TICs</b>	<	3.6e0	<	5.1e0	<	5.1e0		1.1e2		1.8e-6		1.4e-5
Furan, 2,5-dimethyl-	N,J,M		N,J,M	4.4e0	N,J,M		N,J,M		N,J,M		N,J,M	1.2e-5
3-Hexanone	N,J,M	4.3e1	N,J,M	6.0e1	N,J,M	6.1e1	N,J,M	1.2e3	N,J,M	2.1e-5	M,J,M	1.6e-4
2-Hexanone	N,J,M	5.2e1	N,J,M	7.3e1	N,J,M	7.4e1	N,J,M	1.5e3	N,J,M	2.5e-5	N,J,M	2.0e-4
Octane, 3-methyl-	N,J,M	1.8e0	N,J,M	2.5e0	N,J,M	2.5e0	N,J,M	5.2e1	N,J,M	8.6e-7	N,J,M	6.9e-6
Benzaldehyde	N,J,M	2.5e2	N,J,M	3.5e2	N,J,M	3.5e2	N,J,M	7.2e3	N,J,M	1.2e-4	N,J,M	9.6e-4
Dodecane	N,J,M	1.5e1	N,J,M	2.1e1	N,J,M	2.1e1	N,J,M	4.4e2	N,J,M	7.3e-6	N,J,M	5.8e-5
Tridecane	N,J,M	4.5e0	M,L,M	6.3e0	N,J,M	6.4e0	N,J,M	1.3e2	N,J,M	2.2e-6	N,J,M	1.7e-5
Tetradecane	N,J,M	1.9e0	N,J,M	2.7e0	N,J,M	2.8e0	N,J,M	5.6e1	N,J,M	9.4e-7	N,J,M	7.5e-6
Pentadecane	N,J,M	3.4e0	N,J,M	4.7e0	N,J,M	4.8e0	N,J,M	9.8e1	N,J,M	1.6e-6	N,J,M	1.3e-5
Phosphoric acid tributyl ester	N,J,M	5.2e0	N,J,M	7.3e0	N,J,M	7.4e0	M,J,M	1.5e2	N,J,M	2.5e-6	N,J,M	2.0e-5
Cyclododecane	N,J,M	7.0e0	N,J,M	9.8e0	N,J,M	9.9e0	N,J,M	2.0e2	N,J,M	3.4e-6	N,J,M	2.7e-5
Heptadecane	N,J,M	2.0e0	N,J,M	2.8e0	N,J,M	2.8e0	N,J,M	5.8e1	N,J,M	9.7e-7	N,J,M	7.7e-6
Eicosane	N,J,M	1.4e0	N,J,M	1.9e0	N,J,M	2.0e0	N,J,M	4.0e1	N,J,M	6.7e-7	N,J,M	5.3e-6
Hexadecanoic acid	N,J,M	1.2e0	N,J,M	1.7e0	N,J,M	1.7e0	N,J,M	3.5e1	N,J,M	5.9e-7	N,J,M	4.7e-6
Octadecanoic acid	N,J,M	8.1e-1	N,J,M	1.1e0	N,J,M	1.2e0	N,J,M	2.4e1	N,J,M	3.9e-7	N,J,M	3.1e-6
Phosphine oxide, triphenyl-	N,J,M	2.0e0	N,J,M	2.8e0	N,J,M	2.8e0	N,J,M	5.7e1	N,J,M	9.5e-7	N,J,M	7.6e-6
Heneicosane	N,J,M	4.5e-1	N,J,M	6.3e-1	N,J,M	6.4e-1	N,J,M	1.3e1	N,J,M	2.2e-7	N,J,M	1.7e-6
Tetratetracontane	N,J,M	1.5e0	N,J,M	2.1e0	N,J,M	2.1e0	N,J,M	4.3e1	N,J,M	7.2e-7	N,J,M	5.7e-6
Heptane, 2,5-dimethyl-												
Heptane, 2,3-dimethyl-											1	
Benzaldehyde, ethyl-	_											
Octodecane	ı		İ								İ	
Nonacosane	ı											
Hexatriacontane	ı											
Tetracosane					*****							
Tetratriacontane	ı											

Table B-2. SVOC-END-1.

# **SVOC SAMPLING DATA SHEET FOR HLLWE TESTS**

Site:		HLLWE Offgas Tie-in	Sampling	Location:	MA	N-OFG-73	Nozzle No	.:			2-01	Est. DP:	0.14	Est. Tsta	ack, °F:	133
Project:		01-1062-01-0866	Duct ID, i	nches:		12	Nozzle Siz	e, in.:			0.3140	Est. K:		Est. vs, t		25.0
Date:		6/18/2001	Static Pre	essure, in. WG:		-17.5	Pitot No.:	71				Est. DH:		Operator	r(s):	FE/RW
Run No.:		0010-END-1	Est. O2, 9	%:		20.5	Pitot Coeff	f.:				Est. DGM				80
Run Type			Est CO2,			0	Meter Box	No.				Meter Box			Pitot:	
Pbar., in.		25.176	Est. Mois				ΔH≥:					Pretest	0.001	cfm @		in. Hg
Tambient,		70	Impinger			9	Y-factor:	5014	(6.3)		1.0328					pass
DGM Vol.	Goal (m³):	3.00	DGM Vol.	Goal (ft <sup>3</sup> ):		127.080	Min. endin	g DGM vo	l. (π <sup>-</sup> ):		486.849	Post-test	0.000	cfm @	8	in. Hg
Sampling Time	Clock Time	Velocity ΔP	Meter ΔH	Meter Volume	11			MPERATURE	(°F)			Pump	%I,		COMMENT	c
(min.)	(24hr)	(in. WG)	(in. WG)	(cubic feet)	Heated Line	Stack	tn Me	Out	Filter	Impinger Exit	Aux. (XAD)	Vacuum (in. Hg)	764		COMMENT	3
0	15:00	0.14	1.30	359.769	257	132	84	75	262	50	52	7.9		O2=20.5		
10	15:10	0.14	1.30	366.680	257	132	88	76	263	44	51	7.9	104	O2=20.5		
20	15:20	0.14	1.25	373.830	257	131	90	78	262	46	51	7.5	107	O2=20.5		
30	15:30	0.14	1.25	380.845	252	131	91	79	261	47	51	7.5	105	O2=20.5		
40	15:40	0.14	1.25	387.920	255	131	93	80	262	47	52	7.5	105	O2=20.5		
50	15:50	0.14	1.25	395.020	250	131	94	81	262	48	52	7.5	105	O2=20.5		
60	16:00	0.14	1.25	402.150	249	131	94	82	261	48	53	7.5	106	O2=20.5		
70	16:10	0.14	1.25	409.290	249	131	95	82	261	47	51	7.5	106	O2=20.6		
80	16:20	0.14	1.25	416.870	249	131	95	82	261	46	50	7.6	112	O2=20.5		
90	16:30	0.14	1.25	423.600	249	131	95	82	262	46	50	7.5	100	O2=20.6		
100	16:40	0.14	1.25	430.740	249	131	95	83	261	46	50	7.5	106	O2=20.5		
110	16:50	0.14	1.25	437.900	249	131	95	83	262	46	51	7.6	106	O2=20.6		
120	17:00	0.14	1.25	445.075	249	131	95	83	262	46	51	7.5	106	O2=20.5		
130	17:10	0.14	1.25	452.200	249	131	95	83	260	46	52	7.5	105	O2=20.5		
140	17:20	0.14	1.25	459.330	249	131	95	83	261	47	52	7.5	106	O2=20.5		
150	17:30	0.14	1.25	466.570	249	131	95	82	260	47	52	7.5	107	O2=20.5		
160	17:40	0.14	1.25	473.610	249	131	95	82	260	47	51	7.5	104	O2=20.5		
170	17:50	0.14	1.25	480.770	249	131	94	82	260	46	51	7.5	106	O2=20.5		
180	18:00	0.14	1.25	487.903	249	131	94	82	261	46	50	7.5	106	O2=20.6		
Total	Total	ΔPavg	Average	Total		1	Average	Temperat	ures (°F)			Max.	Ave. %li	<u> </u>	-	
180	3:00	0.140	1.26	128.134	251	131	93	81	261	47	51	7.9	106	Avg. O2=2	0.5	

Table B-2. SVOC-END-1.

# 0010 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site:	HLLWE Offg	as Tie-in	1	mpinger Box no.:		9		
Date:	6/18/2001		>	KAD trap Quanterr	ra No.:	A-3405		
Run No.:	0010-END-1							7
Component:	XAD	KO-1	Imp-1	Imp-2		Acid Scrub Section	<u> </u>	_
Туре:	trap	short stem	modified	G-S	short stem	modified	modified	
Reagent:	XAD-2	None	Organics for	ree water	None	2N NaOH	Silica Gel	
Nominal Contents:	20 - 40g, dry	Empty	100 mL	100 mL	Empty	100 mL	300-400g	
Post-test Wt., g:	301.4	534.9	686.9	691.5	574.2	731.0	802.8	Impinger Wt. Gain
Pre-test Wt., g:	300.7	532.0	687.8	691.1	572.8	730.7	<b>78</b> 1.2	Impinger W. Gain
Wt. Gain, g:	0.7	2.9	-0.9	0.4	1.4	0.3	21.6	26.4
Post-test Volume:		0.0	100.0	100.0	0.0			Impinger Vol. Gain
Pre-test Volume:		0.0	100.0	100.0	0.0			mpinger ven eum
Volume Gain:		0.0	0.0	0.0	0.0			0.0
Post-test pH:			6.0	6.0		14.0		
	Filter Lot #	STL-A4023	OF water Lot #	QCLAB-1	NaOH Lot#	QCLAB-381		
O2%	6 20.6	ALLANASTE PET AT THE			_			
CO29	% O.0	_						

Record impinger change-out and other important information below:

Table B-2. SVOC-END-1.

Project:	01-1062-0		
Run Date:			
Run Identification:			
PARAMETER	SYMBOL		22.222
Absolute Pressure in the Duct	Pabs	in. Hg	23.889
Average Duct Gas Temperature	Ts	R	591
Average Meter Temperature	Tm	R	547
Average Gas Oxygen Content	Co2,m	%	20.6
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0
Total Impinger Weight Gain (water)	Ww	grams	26.4
Nozzle Area	An	ft²	0.000538
Duct Area	As	ft²	0.7854
Sample Volume	VmStd	dscf	107.847
Sample Volume (SI)	VmStdm	dscm	3.054
Average Sampling Rate	Qm	dscf/m	0.599
Volume of Water Vapor	VwStd	scf	1.245
Volume of Water Vapor (SI)	VwStdm	scm	0.0352
Moisture Fraction	Bws	-	0.011
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.70
Gas Velocity at Nozzle	vn	ft/s	24.9
Gas Velocity at Nozzle (SI)	vnm	m/s	7.60
Average Gas Velocity	vncor	ft/s	21.05
Dry Offgas Flow Rate	Qsd	dscf/h	41,960
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,188.2
Actual Offgas Flow Rate	Q	acf/h	59,517
Intermediate Isokinetic Rate	li	%	105.8
Final Isokinetic Rate	I	%	105.6

Table B-2. SVOC-END-1.

Project: 01-1062-010866
Run Date: 6/18/2001
Run Identification: 0010-END-1
Run Type:
Lab Report Date: 8/28/2001
Lab Report Status: (preliminary or final)
Final

fina	l)	rinai										
				CONCENTRATIONS				MASS FLOW RATES				
		Actual		Standard (ug/scm)	1	ry Standard (µg/dscm)		μg/min		grams/sec		lb/h
A		ug/acm)	<	(µg/scm) 3.2e0	~	3.2e0	<	6.4e1	_	1.1e-6	<	8.4e-6
Acenaphtheles	< <	2.3e0 2.2e0		3.2e0 3.1e0	ı ·	3.2e0 3.1e0		6.2e1	<	1.0e-6		8.1e-6
Acenaphthylene Acetophenone	<,J	8.3e0			,J	1.2e1		2.3e2		3.9e-6		3.1e-5
Aniline	<	2.5e1	<,5	3.6e1	< "	3.6e1	<,0	7.1e2		1.2e-5		9.4e-8
Anthracene	<	2.2e0	<	3.1e0	<	3.1e0	<	6.2e1	<	1.0e-6		8.1e-6
Benzidine	<	1.5e2	<	2.2e2	<	2.2e2	<	4.3e3	<	7.2e-5	<	5.7e-4
Benzoic acid	<,E	6.0e2	<,E	8.4e2	<,E	8.5e2	<,E	1.7e4	<,E	2.8e-4	<,E	2.2e-3
Benzo(a)anthracene	<	2.8e0	<	3.9e0	<	3.9e0	<	7.8e1	<	1.3e-6	<	1.0e-5
Benzo(a)pyrene	<	3.0e1	<	4.2e1	<	4.3e1	<	8.4e2		1.4e-5		1.1e-4
Benzo(b)fluoranthene	<	6.9e1	<	9.7e1	<	9.8e1	<	1.9e3		3.2e-5		2.6e-4
Benzo(g,h,i)perylene	<	3.9e1	<	5.5e1	<	5.6e1	<	1.1e3	<	1.8e-5	<	1.5e-4
Benzo(k)fluoranthene	<	9.9e1	<	1.4e2	< .	1.4e2	<	2.8e3	<	4.6e-5	<	3.7e-4
Benzyl alcohol	<	1.3e2	<	1.8e2	<	1.9e2	<	3.7e3		6.2e-5	<	4.9e-4
bis(2-Chloroethoxy)methane	<	2.3e0		3.2e0	<	3.3e0	<	6.5e1	<	1.1e-6		8.6e-6
bis(2-Chloroethyl)ether	<	2.5e0		3.6e0	<	3.6e0	<	7.1e1		1.2e-6	ı	9.4e-6 1.9e-4
bis(2-Ethylhexyl)phthalate	< <	5.1e1	< <	7.1e1 3.0e0	<	7.2e1 3.1e0	-	1.4e3 6.1e1	<	2.4e-5 1.0e-6	<	8.1e-6
4-Bromophenyl-phenylether	<	2.2e0 3.0e0	<	4.2e0	2	4.3e0	>	8.4e1	<	1.4e-6	2	1.1e-5
Butylbenzylphthalate	<	3.0e0	<	4.2e0 4.2e0	[	4.3e0 4.3e0	>	8.4e1	<	1.4e-6	<	1.1e-5
Carbazole	<	3.9e0	<	5.5e0	<	5.6e0	<	1.1e2		1.8e-6	<	1.5e-5
4-Chloro-3-methylphenol 4-Chloroaniline	<	2.1e1	<	2.9e1	   ~	2.9e1	<	5.8e2		9.7e-6	<	7.7e-5
2-Chloronaphthalene	2	2.1e0	<	3.0e0	<	3.0e0	<	6.0e1		1.0e-6	<	8.0e-6
2-Chlorophenol	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6		9.4e-6
4-Chlorophenyl phenyl ether	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6	<	9.4e-€
Chrysene	<	2.8e0		3.9e0	<	3.9e0	<	7.8e1	<	1.3e-6	<	1.0e-5
Di-n-butylphthalate	<,J	2.3e1	<,J	3.2e1	<,J	3.3e1	<,J	6.5e2	<,J	1.1e-5	<,J	8.6e-5
Di-n-octylphthalate	<,J	3.7e1	<,J	5.2e1	<,J	5.2e1	<,J	1.0e3		1.7e-5		1.4e-4
Dibenz(a,h)anthracene	<	3.9e1	<_	5.5e1	<	5.6e1	<	1.1 <u>e3</u>		1.8e-5	<	1.5e-4
Dibenzofuran	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6	<	9.4e-6
1,2-Dichlorobenzene	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1		1.2e-6		9.4e-6
1,3-Dichlorobenzene	<b> </b> <	2.8e0	<	3.9e0	<	3.9e0	<	7.8e1		1.3e-6		1.0e-5
1,4-Dichlorobenzene	_ J	4.4e0		6.2e0	J	6.2e0 3.6e1	J <	1.2e2 7.1e2		2.1e-6 1.2e-5		1.6e-5 9.4e-5
3,3'-Dichlorobenzidine	< <	2.5e1 3.0e0		3.6e1 4.2e0	<	4.3e0	<	7.1e2 8.4e1	<	1.4e-6		1.1e-5
2,4-Dichlorophenol	<	3.5e0	<	4.2e0 4.9e0	<	4.9e0	~	9.7e1	<	1.6e-6		1.3e-5
Diethylphthalate	<	2.2e0	<	3.1e0	<	3.1e0	<	6.2e1	<	1.0e-6		8.2e-6
Dimethyl phthalate 2.4-Dimethylphenol	\ <u>`</u>	1.4e1	<	1.9e1	<	1.9e1	<	3.8e2	_	6.4e-6	<	5.1e-5
4,6-Dinitro-2-methylphenol	<	3.0e1		4.2e1	<	4.3e1	<	8.4e2		1.4e-5	<	1.1e-4
2.4-Dinitrophenol	<	6.2e1		8.7e1	<	8.8e1	<	1.8e3		2.9e-5	<	2.3e-4
2.4-Dinitrotoluene	<	3.0e0	<	4.2e0	<	4.3e0	<	8.4e1	<	1.4e-6		1.1e-5
2,6-Dinitrotoluene	<	2.8e0	<	3.9e0	<	3.9e0	<	7.8e1		1.3e-6		1.0e-5
1,2-Diphenylhydrazine	<	2.3e0	<	3.2e0	<	3.2e0	<	6.4e1		1.1e-6		8.4e-6
Fluoranthene	<	2.3e0	<	3.2e0	<	3.3e0	<	6.5e1		1.1e-6		8.6e-6
Fluorene	<	2.3e0	<	3.2e0	<	3.3e0	<	6.5e1	<	1.1e-6		8.6e-6
Hexachlorocyclopentadiene	<	3.7e1	<	5.2e1	<	5.2e1	<	1.0e3		1.7e-5	<	1.4e-4
Hexachlorobenzene	<	2.5e0		3.6e0	<	3.6e0	< <	7.1e1	<	1.2e-6		9.4e-6
Hexachlorobutadiene	<	3.5e0	<	4.9e0	< <	4.9e0	<	9.7e1 1.0e2	<	1.6e-6 1.7e-6	<	1.3e-5 1.4e-5
Hexachloroethane	< <	3.7e0 3.5e1	<	5.2e0 4.9e1	<	5.2e0 4.9e1	~	9.7e2	<	1.7e-0	<	1.3e-4
Indeno(1,2,3-cd)pyrene	\ <u></u>	2.3e0	,	4.9e1 3.2e0	<	3.3e0	<	6.5e1	<	1.1e-6	<	8.6e-6
Isophorone	<	2.3e0 2.3e0		3.2e0	<	3.3e0	~	6.5e1	<	1.1e-6		8.6e-6
2-Methylnaphthalene 2-Methylphenol	<	1.1e1	<	1.5e1		1.5e1	<	3.0e2		5.1e-6	ı	4.0e-5
3-Methylphenol & 4-Methylphenol	<	7.8e0		1.1e1	<	1.1e1	<	2.2e2		3.7e-6		2.9e-5
N-Nitroso-di-n-propylamine	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	ı	1.2e-6		9.4e-6
N-Nitrosodimethylamine	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6	<	9.4e-€
N-Nitrosodiphenylamine	<	3.5e0	<	4.9e0	<	4.9e0	<	9.7e1	<	1.6e-6	< .	1.3e-5
Naphthalene	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6	<	9.4e-6
2-Nitroaniline	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6	<	9.4e-6
3-Nitroaniline	<	9.5e0	<	1.3e1	<	1.3e1	<	2.7e2		4.4e-6	<	3.5e-6
4-Nitroaniline	<	8.3e0	<	1.2e1	<	1.2e1	<	2.3e2		3.9e-6		3.1e-5
Nitrobenzene	<,J	3.5e0	<,J	4.9e0	<b>&lt;</b> ,J	4.9e0	<,J	9.7e1		1.6e-6		1.3e-5
2-Nitrophenol	<,J	9.5e0		1 3e1	<.J	1 3e1	<,J	2.7e2	1	4.4e-6		3.5e-5
4-Nitrophenol	<	1.3e1	<		<	1.8e1	<	3.6e2		5.9e-6		4.7e-5
2,2'-Oxybis(1-chloropropane)	<	3.2e0	<	4.5e0	<	4.6e0	<b>!</b> <	9.1e1	I <sup>&lt;</sup> .	1.5e-6	l <u></u>	1.2e-5

Table B-2. SVOC-END-1.

hriai			CONC	ENTRATIO	TRATIONS MASS FLOW RATES							
	Ac	tual		tandard		tandard					1	
	(μg/	acm)	()	ıg/scm)	(µg/	dscm)	µg/	min .	gran	ns/sec	- 1	b/h
Pentachlorobenzene	<	2.3e0	<	3.2e0	<	3.3e0	<	6.5e1	<	1.1e-6	<	8.6e-6
Pentachloronitrobenzene	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6	<	9.4e-6
Pentachiorophenol	<	6.9e1	<	9.7e1	<	9.8e1		1.9e3		3.2e-5		2.6e-4
Phenanthrene	<	2.3e0	<	3.2e0	<	3.2e0		6.4e1		1.1e-6		8.4e-6
Phenol	<,J	8.5e0	<,J	1.2e1	<,J	1.2e1	<,J	2.4e2	<,J	4.0e-6	<,J	3.2e-5
Pyrene	<	2.3e0		3.2e0	<	3.3e0		6.5e1		1.1e-6		8.6e-6
Pyridine	<	3.9e0	<	5.5e0	<	5.6e0		1.1e2		1.8e-6		1.5e-5
1,2,4,5-Tetrachlorobenzene	<	2.5e0	<	3.6e0	<	3.6e0	<	7.1e1	<	1.2e-6		9.4e-6
1,2,4-Trichlorobenzene	<	2.8e0		3.9e0		3.9e0		7.8e1		1.3e-6		1.0e-5
2,4,5-Trichlorophenol	<	5.8e0	<	8.1e0	<	8.2e0		1.6e2		2.7e-6		2.1e-5
2,4,6-Trichlorophenol	<	3.7e0	<	5.2e0	<	5.2e0	<	1.0e2	<	1.7e-6	<	1.4e-5
TICs			l									
Furan, 2,5-dimethyl-	N,J,M	1.8e0	N,J,M	2.5e0	N,J,M	2.6e0	N,J,M		N,J,M		N,J,M	6.7e-6
3-Hexanone	N,J,M		N,J,M		N,J,M		N,J,M		N,J,M		N,J,M	8.2e-5
2-Hexanone	N,J,M	2.8e1	N,J,M		N,J,M		N,J,M		N,J,M		N,J,M	1.0e-4
Heptane, 2,3-dimethyl-	N,J,M		N,J,M	3.2e0	N,J,M		N,J,M		N,J,M		N,J,M	8.5e-6
Benzaldehyde	N,J,M	1.7e2	N,J,M	2.4e2	N,J,M		N,J,M		N,J,M		N,J,M	6.3e-4
Formic acid, phenylmethyl ester	N,J,M	1.7e1	N,J,M		N,J,M		N,J,M		N,J,M		N,J,M	6.2e-5
Benzaldehyde, 4-ethyl-	N,J,M	1.3e1	N,J,M		N,J,M		N,J,M		N,J,M		N,J,M	4.9e-5
Dodecane	N, i, M		M,J,M		N,J,M		N,J,M		N,J,M		N,J,M	4.1e-5
Tridecane	N,J,M		M,J,M		N,J,M		N,J,M		N,J,M		N,J,M	1.8e-5
2,4-Hexadiene	N,J,M		M,J,M		N,J,M		N,J,M		N,J,M		N,J,M	1.4e-4
Tetradecane	N,J,M		M,L,N		N,J,M		N,J,M		M,J,M		N,J,M	8.2e-5
Phosphoric acid tributyl ester	N,J,M		N,J,M		N,J,M		N,J,M		N,J,M		N,J,M	1.4e-5
Cyclododecane	N,J,M		N,J,M		N,J,M		N,J,M		N,J,M		N,J,M	2.6e-5
Heptadecane	N,J,M	9.2e-1			N,J,M		N,J,M		N,J,M		N,J,M	3.4e-6
Octadecanoic acid	N,J,M	4.6e-1	N,J,M	6.5e-1	N,J,M	6.5e-1	N,J,M	1.3e1	N,J,M	2.2e-7	N,J,M	1.7e-6
Heptane, 2,5-dimethyl-												
Benzoic acid, methyl ester					1							
Benzaldehyde, ethyl-	ŀ											
Pentadecane												
Eicosane									ļ		ļ	
Heneicosane												
Octodecane												
Phosphine oxide, triphenyl-												
Nonacosane												
Hexatriacontane												
Tetracosane												
Tetratriacontane	ļ		L						Ц		1	

Table B-3. SVOC-STRT-2.

# **SVOC SAMPLING DATA SHEET FOR HLLWE TESTS**

Site:		HLLWE Offgas Tie-in	Sampling	Location:	1AM	N-OFG-73	Nozzle No	:			2-01	Est. DP:	0.16	Est. Tstack	k, °F:	133
Project:		01-1062-01-0866	Duct ID, i	nches:		12	Nozzle Siz	e, in.:			0.3140	Est. K:	6.53	Est. vs, ft/s	s:	26.8
Date:		6/19/2001	Static Pre	ssure, in. WG:		-17.5	Pitot No.:					Est. DH:		Operator(s	s):	FE/RW
Run No.:		0010-STRT-2				20.5	Pitot Coeff	:			0.84	Est. DGM	Temperatur	re, °F		
Run Type:			Est CO2,			0	Meter Box	No.					Leak Che	cks:	Pitot:	pass
Pbar., in. I			Est. Moist			1.3%				<del></del>		Pretest	0.003	cfm @		in. Hg
Tambient,			Impinger			4	Y-factor:		3.		1.0328					pass
DGM Vol.	Goal (m³):	3.00	DGM VOI.	Goal (ft <sup>3</sup> ):		127.080	Min. endin				615.228	Post-test	0.000	cfm @	6 i	in. Hg
Sampling	Clock	Velocity ΔP	Meter ΔH	Meter				PERATURE	(°F)			Pump	%l,		O MENTO	
Time (min.)	Time (24hr)	(in. WG)	(in. WG)	Volume (cubic feet)	Heated Line	Stack	In Me	ter Out	Filter	lmpinger Exit	Aux. (XAD)	Vacuum (in. Hg)	701;	١	COMMENTS	
0	8:00	0.16	1.30	488.148	267	132	62	53	262	46	48	6.0	-			
10	8:10	0.16	1.30	494.990	270	133	69	53	263	43	46	6.0	100			
20	8:20	0.16	1.30	501.989	273	133	73	59	267	44	47	6.0	101			
30	8:30	0.16	1.30	508.995	273	132	75	61	262	44	47	6.0	101			
40	8:40	0.16	1.30	515.900	271	132	77	63	269	45	47	6.0	99			
50	8:50	0.16	1.30	523.020	271	132	79	65	263	45	47	6.0	102			
60	9:00	0.16	1.30	530.080	270	132	80	66	264	46	48	6.0	101			
70	9:10	0.16	1.30	537.101	271	132	81	68	264	46	48	6.0	100			
80	9:20	0.16	1.30	544.260	270	132	82	69	262	47	48	6.0	102			
90	9:30	0.16	1.30	551.510	271	132	83	70	262	47	48	6.0	103			
100	9:40	0.16	1.30	558.270	272	132	84	71	262	48	50	6.0	96			
110	9:50	0.16	1.30	565.420	271	132	85	72	263	48	50	6.0	101			
120	10:00	0.16	1.30	572.570	271	132	85	72	263	48	51	6.0	101			
130	10:10	0.16	1.30	579.710	269	132	86	73	263	49	53	6.0	101			
140	10:20	0.16	1.30	586.860	270	132	87	74	263	49	53	6.0	100			
150	10:30	0.16	1.30	593.900	271	132	87	74	264	50	56	6.0	99			
160	10:40	0.16	1.30	601.150	269	132	88	75	264	50	57	6.0	102			
170	10:50	0.16	1.30	608.300	271	132	88	76	262	50	59	6.0	100			
180	11:00	0.16	1.30	615.462	270	132	89	76	262	51	61	6.0	100			
Total	Total	ΔPavg	Average	Total			Average	Temperat	1	1	<del></del>	Max.	Ave. %I	1		
180	3:00	0.160	1.30	127.314	271	132	81	68	263	47	51	6.0	100			

Table B-3. SVOC-STRT-2.

# 0010 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site:	HLLWE O	ffgas Tie-in		Impinger Box no.:	4	_		
Date:	6/19/2001			XAD trap Quanterr	A-3374	_		
Run No.:	0010-STRT-2							
Component:	XAD	KO-1	Imp-1	lmp-2		Acid Scrub Section	n	
Туре:	trap	short stem	modified	G-S	short stem	modified	modified	
Reagent:	XAD-2	None	Organics	free water	None	2N NaOH	Silica Gel	
Nominal Contents:	20 - 40g, dry	Empty	100 mL	100 mL	Empty	<b>100</b> mL	300-400g	1
Post-test Wt., g:	288.7	548.8	696.5	683.4	536.1	730.9	837.6	Impinger Wt.
Pre-test Wt., g:	288.5	546.3	698.3	683.7	535.0	733.5	813.2	Gain
Wt. Gain, g:	0.2	2.5	-1.8	-0.3	1.1	-2.6	24.4	23.5
Post-test Volume:		0.0	100.0	100.0	0.0			Impinger Vol.
Pre-test Volume:		0.0	100.0	100.0	0.0			Gain
Volume Gain:		0.0	0.0	0.0	0.0			0.0
Post-test pH:			6.0	6.0		13.0		
	Filter Lot #	STL-A4023	OF water Lot #	QCLAB-1	NaOH Lot#	000381		_
O2%	20.6	· · · · · · · · · · · · · · · · · · ·	_		-		-	
CO2%	0.0							

Record impinger change-out and other important information below:

Table B-3. SVOC-STRT-2.

Project: 01-1062-01-0866									
Run Date:									
Run Identification:	0010-S SYMBOL		<del></del>						
PARAMETER Absolute Pressure in the Duct	Pabs	UNITS in. Hg	23.805						
	, 5								
Average Duct Gas Temperature	Ts	R	592						
Average Meter Temperature	Tm	R	534						
Average Gas Oxygen Content	Co2,m	%	20.6						
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0						
Total Impinger Weight Gain (water)	Ww	grams	23.5						
Nozzle Area	An	ft²	0.000538						
Duct Area	As	ft²	0.7854						
Sample Volume	VmStd	dscf	109.351						
Sample Volume (SI)	VmStdm	dscm	3.096						
Average Sampling Rate	Qm	dscf/m	0.608						
Volume of Water Vapor	VwStd	scf	1.108						
Volume of Water Vapor (SI)	VwStdm	scm	0.0314						
Moisture Fraction	Bws	-	0.010						
Dry Gas Molecular Weight	Md	g/g-mol	28.82						
Wet Gas Molecular Weight	Ms	g/g-mol	28.72						
Gas Velocity at Nozzle	vn	ft/s	26.7						
Gas Velocity at Nozzle (SI)	vnm	m/s	8.15						
Average Gas Velocity	vncor	ft/s	22.56						
Dry Offgas Flow Rate	Qsd	dscf/h	44,791						
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,268.3						
Actual Offgas Flow Rate	Q	acf/h	63,775						
Intermediate Isokinetic Rate	li	%	100.6						
Final Isokinetic Rate	ı	%	100.3						

### Table B-3. SVOC-STRT-2.

Project: 01-1062-01-0866
Run Date: 6/19/2001
Run Identification: 0010-STRT-2
Run Type: Test
Lab Report Date: 8/28/2001
Lab Report Status: (preliminary or final)
Final

### **RESULTS**

	al)		ON	CENTRATION	ONS MASS FLOW RATES									
	-	Actual	I	Standard		ry Standard								
		(µg/acm)		(µg/scm)		(µg/dscm)		μg/min		grams/sec		lb/h		
Acenaphthene	<,J	3.6e0	<	5.1e0		5.2e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-5		
Acenaphthylene	<,J	3.6e0		5.1e0		5.2e0		1.1e2		1.8e-6		1.4e-		
Acetophenone	-,J	9.5e0		1.3e1		1.4e1	<,J	2.9e2		4.8e-6		3.8e-5		
Aniline	<'-	3.4e1		4.8e1	<	4.8e1	<	1.0e3		1.7e-5	<	1.4e-4		
Anthracene	<,J	3.6e0		5.1e0	<.J	5.2e0	<	1.1e2		1.8e-6	<,J	1.4e-5		
Benzidine	< '	1.8e2		2.6e2	< '-	2.6e2		5.5e3		9.1e-5	<'-	7.2e-4		
Benzoic acid	lΕ	5.2e2		7.4e2	F	7.4e2	1		E	2.6e-4	E	2.1e-3		
Benzo(a)anthracene	<,,	4.3e0		6.1e0		6.1e0		1.3e2		2.2e-6	_	1.7e-5		
Benzo(a)pyrene	<,J	3.2e1	<,J	4.5e1	<,J	4.5e1	<,J	9.6e2	<,J	1.6e-5	<,J	1.3e-4		
Benzo(b)fluoranthene	,, <,J	6.8e1		9.6e1		9.7e1	<,J	2.0e3	<,J	3.4e-5	<,J	2.7e-4		
Benzo(g,h,i)perylene	<,J	4.1e1		5.8e1		5.8e1		1.2e3		2.0e-5	<,J	1.6e-4		
Benzo(k)fluoranthene	<,J	9.8e1	<,J	1.4e2		1.4e2	<,J	2.9e3	<,J	4.9e-5	<,J	3.9e-4		
Benzyl alcohol	<	1.3e2		1.8e2	<	1.8e2	<	3.9e3	<	6.5e-5	<	5.1e-4		
bis(2-Chloroethoxy)methane	<	3.2e0		4.5e0	<	4.5e0	<	9.6e1	<	1.6e-6	<	1.3e-5		
bis(2-Chloroethyl)ether	<.J	3.6e0		5.1e0		5.2e0	1	1.1e2	<,J	1.8e-6	ر,>	1.4e-5		
bis(2-Ethylhexyl)phthalate	<,J	2.9e1	<,J	4.2e1	<,J	4.2e1	<,J	8.9e2	<,J	1.5e-5	< ,	1.2e-4		
4-Bromophenyl-phenylether	<,J	4.1e0		5.8e0		5.8e0	<,J	1.2e2	<,J	2.0e-6	<,J	1.6e-5		
Butylbenzylphthalate	<,J	4.1e0 4.3e0		6.1e0		6.1e0		1.3e2		2.2e-6		1.7e-5		
Carbazole	<,J	4.1e0		5.8e0		5.8e0	<,J	1.2e2		2.0e-6		1.6e-5		
4-Chloro-3-methylphenol	<,J	7.0e0		9.9e0		1.0e1	<,J	2.1e2	<,J	3.5e-6	<,J	2.8e-5		
4-Chloroaniline	<	2.5e1		3.5e1	<,0	3.6e1	<	7.5e2	<	1.3e-5	<	9.9e-5		
2-Chloronaphthalene	\ \ \ \	3.4e0		4.8e0		4.8e0			-  <,J	1.7e-6		1.4e-5		
2-Chlorophenol	د, <i>&gt;</i> ال, >			4.8e0		4.8e0	<,J	1.0e2		1.7e-6	>,J <,J	1.4e-5		
	<,J	3.4e0 3.9e0	<,J <,J	5.4e0		5.5e0	<,J	1.0e2 1.2e2	<b>&gt;</b> ,J	1.7e-6 1.9e-6	J	1.5e-5		
4-Chlorophenyl phenyl ether	<,J	4.8e0	<,J	6.7e0		6.8e0		1.4e2		2.4e-6		1.9e-5		
Chrysene		2.5e1		3.5e1		3.6e1		7.5e2	-,J  -,J	1.3e-5	<.J	9.9e-5		
Di-n-butylphthalate	<,J	2.5e1 3.6e1	<,j	5.1e1		5.2e1	<,J	7.5e2 1.1e3		1.8e-5	ر, د ار >	1.4e-4		
Di-n-octylphthalate	<,J		<,J						[<,J					
Dibenz(a,h)anthracene	<,J	3.9e1	<,J	5.4e1		5.5e1	<,J	1.2e3	<u>&lt;,J</u>	1.9e-5	<,J	1.5e-4		
Dibenzofuran	<,J	3.9e0	<,J	5.4e0		5.5e0	<,J	1.2e2	<,J	1.9e-6	<,J	1.5e-5		
1,2-Dichlorobenzene	<,J	3.6e0				5.2e0	<,J	1.1e2	<,J	1.8e-6		1.4e-5		
1,3-Dichlorobenzene	<,J	3.9e0		5.4e0	<,J	5.5e0	•		<b> </b> <,J	1.9e-6		1.5e-5		
1,4-Dichlorobenzene	J	4.5e0	J	6.4e0	J	6.5e0	J	1.4e2	J.	2.3e-6	J	1.8e-5		
3,3'-Dichlorobenzidine	<	2.9e1	<	4.2e1		4.2e1	<	8.9e2	<	1.5e-5	<	1.2e-4		
2,4-Dichlorophenol	<b> </b> < .	4.1e0	< .	5.8e0		5.8e0	<b> </b> <	1.2e2	۲.	2.0e-6	< .	1.6e-5		
Diethylphthalate	ر,>	5.0e0		7.0e0		7.1e0		1.5e2	<,J	2.5e-6		2.0e-5		
Dimethyl phthalate	<,J	3.6e0	<u>&lt;,J</u>	5.1e0	<,J	5.2e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-5		
2,4-Dimethylphenol	<	1.4e1	<		<	2.0e1	<	4.2e2	<	7.1e-6	<	5.6e-5		
4,6-Dinitro-2-methylphenol	<		<	4.5e1		4.5e1	<	9.6e2	<	1.6e-5	<	1.3e-4		
2,4-Dinitrophenol	<	6.4e1	<	9.0e1		9.0e1	<	1.9e3	<	3.2e-5	<b> </b> < .	2.5e-4		
2,4-Dinitrotoluene	<,J	4.3e0		6.1e0		6.1e0		1.3e2	<,J	2.2e-6	<,J	1.7e-5		
2,6-Dinitrotoluene	<,J			5.1e0		5.2e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-5		
1,2-Diphenylhydrazine	<,J	3.6e0	<,J	5.1e0		5.2e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-5		
Fluoranthene	<,J		<,J		<,J	5.5e0	<,J	1.2e2	<,J	1.9e-6	<,J	1.5e-5		
Fluorene	<,J		<,J	5.1e0		5.2e0	<,J	1.1e2	<b>&lt;</b> ,J	1.8e-6	<,J	1.4e-5		
Hexachlorocyclopentadiene	<	4.1e1	<	5.8e1	<	5.8e1	<	1.2e3	<	2.0e-5	<	1.6e-4		
Hexachlorobenzene	<,J		<,J	5.1e0		5.2e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-5		
Hexachlorobutadiene	ال,>	4.5e0	<,J		<,J	6.5e0	<,J	1.4e2	<,J	2.3e-6	<,J	1.8e-5		
Hexachloroethane	<,J	4.5e0	<,J		<,J	6.5e0	<,J	1.4e2	<,J	2.3e-6	<,J	1.8e-5		
Indeno(1,2,3-cd)pyrene	<,J	3.4e1	<,J	4.8e1	<,J	4.8e1	<,J	1.0e3	<,J	1.7e-5	<,J	1.4e-4		
Isophorone	ر,>	3.6e0	<,J		۷,>	5.2e0	<,J	1.1e2	<,J	1.8e-6	<,J	1.4e-5		
2-Methylnaphthalene	<,J	3.4e0	۷,>		<,J	4.8e0			<,J	1.7e-6	<,J	1.4e-5		
2-Methylphenol	<	1.2e1	<	1.6e1	<	1.6e1	<	3.5e2	<	5. <b>8e-</b> 6	<	4.6e-5		
3-Methylphenol & 4-Methylphenol	<	8.8e0	<	1.2e1	<	1.3e1	٧		<	4.4e-6	<	3.5e-5		
N-Nitroso-di-n-propylamine	<	3.4e0	<	4.8e0	<	4.8e0	<	1.0e2	<	1.7e-6	<	1.4e-5		
N-Nitrosodimethylamine	<,J	3.4e0	<,J	4.8e0	<,J	4.8e0	<,J	1.0e2	<,J	1.7e-6	ل,>	1.4e-5		
N-Nitrosodiphenylamine	<b>ا</b> ر>	4.8e0	<,J	6.7e0	<,J	6.8e0	<,J	1.4e2	<,J	2.4e-6	<,J	1.9e-5		
Naphthalene	<,J	3.9e0	<,J	5.4e0	<,J			1.2e2	<,J	1.9e-6	<,J	1.5e-5		
2-Nitroaniline	<	3.9e0	<		<	5.5e0	<	1.2e2	<	1.9e-6	<	1.5e-5		
3-Nitroaniline	<	1.1e1	<	1.6e1	<	1.6e1	<		<	5.7e-6	<	4.5e-5		
4-Nitroaniline	<	1.0e1	<		<	1.4e1	<	3.0e2		5.0e-6		4.0e-5		

Table B-3. SVOC-STRT-2.

Project: 01-1062-01-0866 Project: 01-1062-01-0866

Run Date: 6/19/2001

Run Identification: 0010-STRT-2

Run Type: Test

Lab Report Date: 8/28/2001

Lab Report Status: (preliminary or final)

Final

**RESULTS** 

final	)		l .									
			ONCENT				MASS FLOW RATES					
	Actual		Stan	dard		andard	l					
	(µg/acm	1)	(µg/s	scm)	(μg/c	dscm)	μg/	min .	gran	ns/sec	I	b/h
Nitrobenzene	<,J	4.5e0	<,J	6.4e0	<,J	6.5e0	<,J	1.4e2	<,J	2.3e-6	<,J	1.8e-5
2-Nitrophenol		8.4e0	<,J	1.2e1	<,J	1.2e1	<,J	2.5e2	<,J	4.2e-6	<,J	3.3e-5
4-Nitrophenol		1.4e1	<	2.0e1	<	2.0e1	<	4.3e2	<	7.2e-6	<	5.7e-5
2,2'-Oxybis(1-chloropropane)	<,J	5.0e0	<,J	7.0e0	<,J	7.1e0	<,J	1.5e2	< J	2.5e-6	<,J	2.0e-5
Pentachlorobenzene	<	3.4e0	<	4.8e0	<	4.8e0	<	1.0e2	<	1.7e-6	<	1.4e-5
Pentachloronitrobenzene	<	3.6e0	<	5.1e0	<	5.2e0	<	1.1e2	<	1.8e-6	<	1.4e-5
Pentachlorophenol	<	7.0e1	<	9.9e1	<	1.0e2	<	2.1e3	<	3.5e-5	<	2.8e-4
Phenanthrene	<,J	3.9e0	<,J	5.4e0	<,J	5.5e0	<,J	1.2e2	<,J	1.9e-6	<,J	1.5e-5
Phenol	<,J	7.5e0	<,J	1.1e1	<,J	1.1e1	<,J	2.3e2	<,J	3.8e-6	<,J	3.0e-5
Pyrene	<,J	4.1e0	<,J	5.8e0	<,J	5.8e0	<,J	1.2e2	<,J	2.0e-6	<,J	1.6e-5
Pyridine	<	6.4e0	<	9.0e0	<	9.0e0	<	1.9e2	<	3.2e-6	<	2.5e-5
1,2,4,5-Tetrachlorobenzene	<	3.6e0	<	5.1e0	<	5.2e0		1.1e2		1.8e-6		1.4e-5
1,2,4-Trichlorobenzene	<,J	3.9e0	<,J	5.4e0	<,J	5.5e0	<,J	1.2e2	<,J	1.9e-6	<,J	1.5e-5
2,4,5-Trichlorophenol	<	6.6e0	<	9.3e0	<	9.4e0	<	2.0e2	<	3.3e-6		2.6e-5
2,4,6-Trichlorophenol	<	4.8e0	<	6.7e0	<	6.8e0	<	1.4e2	<	2.4e-6	<	1.9e-5
TICs												
Furan, 2,5-dimethyl-	N,J,M	2.2e0	N,J,M	3.1e0	N,J,M	3.2e0	N,J,M		N,J,M		N,J,M	8.8e-6
3-Hexanone	N,J,M	2.2e1	N,J,M	3.1e1	M,J,M		N,J,M		N,J,M	1.1e-5		8.7e-5
Heptane, 2,5-dimethyl-	N,J,M	1.6e0	N,J,M	2.3e0	N,J,M		N,J,M		N,J,M	8.1e-7		6.4e-6
Benzaldehyde	N,J,M	1.5e2	N,J,M		N,J,M		N,J,M		N,J,M	7.6e-5		6.1e-4
Formic acid, phenylmethyl este	N,J,M	1.2e1	N,J,M	1.7e1	N,J,M		N,J,M		N,J,M	5.9e-6		4.7e-5
Dodecane		1.2e1	N,J,M		N,J,M		N,J,M		N,J,M	6.3e-6		5.0e-5
Tridecane	N,J,M	4.1e0	N,J,M	5.8e0	N,J,M		N,J,M		N,J,M	2.0e-6		1.6e-5
Naphthalene, 1-methyl-		1.8e0	N,J,Q		N,J,Q		N,J,Q		N,J,Q	8.9e-7		7.0e-6
Tetradecane			N,J,M	1.8e1	N,J,M		N,J,M		N,J,M	6.4e-6		5.1e-5
Cyclododecane			N,J,M		N,J,M		N,J,M		N,J,M	6.4e-7		5.1e-6
Hexanedioic acid, bis(2-ethylh)			N,J,Q		N,J,Q		N,J,Q		N,J,Q	1.1e-6		9.0e-6
1,2-Benzenedicarboxylic acid,			N,J,Q		N,J,Q		N,J,Q		N,J,Q	9.6e-7		7.6e-6
Benzo(e)pyrene	N,J,Q	3.2e0	D,L,N	4.5e0	D,L,N	4.5e0	N,J,Q	9.6e1	N,J,Q	1.6e-6	N,J,Q	1.3e-5
Heptane, 2,3-dimethyl-												
Benzoic acid, methyl ester												
Benzaldehyde, ethyl-												
Pentadecane	ł											
Heptadecane	ì						İ					
Eicosane	ŀ											
Heneicosane												
Octodecane			İ									
Phosphine oxide, triphenyl-												
Nonacosane	ŀ						1				1	
Hexatriacontane												
Tetracosane											1	
Tetratriacontane			L								L	

Table B-4. SVOC-END-2.

## **SVOC SAMPLING DATA SHEET FOR HLLWE TESTS**

Site:		LILLIANE Office The	C	3V00 3AI						***				I	
Project:		HLLWE Offgas Tie-in	<del> </del>		MA		Nozzie No.					Est. DP:		Est. Tstack, °F:	132
Date:		01-1062-01-0866		ncnes: essure, in. WG:		12 -17.5	Nozzle Siz Pitot No.:	e, in.:			0.3140			Est. vs, ft/s:	25.9
Run No.:		0010-END-2					Pitot No.:					Est. DH:	1.15 Temperatur	Operator(s):	RW,FE
Run Type:			Est CO2,										Leak Che	<del></del>	80 : Pass
Pbar., in. I		25.099	Est. Mois			1.3%		NO.				Pretest			in. Hg
Tambient,		75	Impinger	<del> </del>		2	Y-factor:				1.0328	116656	0.001	Pitot	
DGM vol.		3.00		Goal (ft <sup>3</sup> ):			Min. ending	g DGM vol	l. (ft <sup>3</sup> ):	· · · · · · · · · · · · · · · · · · ·		Post-test	0.001		in. Hg
Sampling	Clock	Velocity	Meter	Meter			TEN	MPERATURE	(°F)			Pump			
Time	Time	ΔΡ	ΔН	Volume	Heated			ter		Impinger	Aux.	Vacuum	%l <sub>i</sub>	COMMENT	's
(min.)	(24hr)	(in. WG)	(in. WG)	(cubic feet)	Line	Stack	In	Out	Filter	Exit	(XAD)	(in. Hg)			
0	14:00	0.14	1.30	619.354	266	132	81	76	275	55	64	12.5	-		
10	14:10	0.14	1.30	626.430	266	132	84	77	276	52	58	12.5	106		
20	14:20	0.14	1.30	633.410	266	132	88	78	259	54	56	12.5	104		
30	14:30	0.14	1.25	640.390	266	132	90	80	260	55	55	12.5	104		
40	14:40	0.14	1.20	647.200	266	132	91	81	259	55	55	12.1	101		
50	14:50	0.14	1.20	654.660	266	132	92	81	260	55	54	12.1	111		
60	15:00	0.14	1.15	661.640	265	132	93	82	260	56	56	11.9	103		
70	15:10	0.14	1.15	668.610	266	132	93	82	260	56	57	11.9	103		
80	15:20	0.14	1.15	675.590	266	132	94	82	260	56	57	11.9	103		***
90	15:30	0.14	1.15	682.560	266	132	94	83	259	<b>5</b> 6	57	11.9	103		
100	15:40	0.14	1.15	689.560	265	132	95	84	259	57	58	11.9	103		
110	15:50	0.14	1.15	696.570	265	132	95	84	259	57	59	11.9	104		
120	16:00	0.14	1.15	703.580	265	132	95	85	259	58	61	11.9	103		
130	16:10	0.14	1.15	710.590	265	132	95	85	259	55	61	11.9	103		
140	16:20	0.14	1.14	717.620	265	132	95	85	258	53	59	11.9	104		
150	16:30	0.14	1.20	724.740	265	132	96	85	259	53	59	11.9	105		
160	16:40	0.14	1.20	731.880	265	132	96	85	258	53	60	11.9	105		
170	16:50	0.14	1.20	738.740	265	132	96	86	259	54	60	11.9	101	_	
180	17:00	0.14	1.20	745.790	265	132	96	86	259	54	61	11.9	104		
190	17:10	0.14	1.20	752.830	265	132	96	85	259	54	61	11.9	104		
195	17:15	0.14	1.20	756.352	265	132	95	86	258	54	62	11.9	104		
Total	Total	ΔPavg	Average	Total			Average	Temperat	ures (°F)			Max.	Ave. %l <sub>i</sub>		
195	3:15	0.140	1.19	<b>13</b> 6.998	265	132	93	83	261	55	59	12.5	104		

Table B-4. SVOC-END-2.

# 0010 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site:	HLLWE Offg	gas Tie-in	1	Impinger Box no.:		2		
Date:	6/19/2001		:	XAD trap Quanter	ra No.:	A-3424	· -	
Run No.:	0010-END-2							_
Component:	XAD	KO-1	lmp-1	Imp-2		Acid Scrub Section	n	
Туре:	trap	short stem	modified	G-S	short stem	modified	modified	
Reagent:	XAD-2	None	Organics f	ree water	None	2N NaOH	Silica Gel	
Nominal Contents:	20 - 40g, dry	Empty	100 mL	100 mL	Empty	100 mL	300-400g	
Post-test Wt., g:	313.3	533.9	681.7	691.6	576.5	728.6	800.7	Impinger Wt. Gain
Pre-test Wt., g:	313.6	531.9	688.2	690.9	573.6	730.4	773.4	Impinger Wt. Gain
Wt. Gain, g:	-0.3	2.0	-6.5	0.7	2.9	-1.8	27.3	24.3
Post-test Volume:		0.0	100.0	100.0	0.0			Impinger Vol. Gain
Pre-test Volume:		0.0	100.0	100.0	0.0			Impinger voi. Gain
Volume Gain:		0.0	0.0	0.0	0.0			0.0
Post-test pH:			6.0	6.0		13.0		
	Filter Lot#	STL-A4023	OF water Lot #	QCLAB-1	NaOH Lot#	000381		
02%	20.6				=		-	
CO2%	0.0							

Record impinger change-out and other important information below:

Table B-4. SVOC-END-2.

Project:		01-0866	
Run Date:	6/19/2		
Run Identification:			
PARAMETER	SYMBOL		
Absolute Pressure in the Duct	Pabs	in. Hg	23.812
Average Duct Gas Temperature	Ts	R	592
Average Meter Temperature	Tm	R	547
Average Gas Oxygen Content	Co2,m	%	20.6
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0
Total Impinger Weight Gain (water)	Ww	grams	24.3
Nozzle Area	An	ft²	0.000538
Duct Area	As	ft²	0.7854
Sample Volume	VmStd	dscf	114.799
Sample Volume (SI)	VmStdm	dscm	3.251
Average Sampling Rate	Qm	dscf/m	0.589
Volume of Water Vapor	VwStd	scf	1.146
Volume of Water Vapor (SI)	VwStdm	scm	0.0324
Moisture Fraction	Bws	-	0.010
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.72
Gas Velocity at Nozzle	vn	ft/s	25.0
Gas Velocity at Nozzle (SI)	vnm	m/s	7.62
Average Gas Velocity	vncor	ft/s	21.09
Dry Offgas Flow Rate	Qsd	dscf/h	41,913
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,186.9
Actual Offgas Flow Rate	Q	acf/h	59,641
Intermediate Isokinetic Rate	li	%	104.2
Final Isokinetic Rate	i .	%	103.9

Table B-4. SVOC-END-2.

Project:

01-1062-01-0866

Run Date: Run Identification: Run Type: Lab Report Date: Lab Report Status: (preliminary or

6/19/2001 0010-END-2 Test 8/28/2001

**RESULTS** 

	T	CO	CENT	RATION	S			MA	SS F	LOW RAT	ES	
	Actua			ndard		Standard						
	(µg/acı	m)	(µg	/scm)	μg	(dscm)	μ	g/min	gra	ms/sec		lb/h
Acenaphthene	<	2.1e0	<	3.0e0	<	3.0e0	<	6.0e1	<	9.9e-7	<	7.9e-
Acenaphthylene	<	2.1e0	<	2.9e0	<	2.9e0	<	5.8e1	<	9.6e-7	<	7.6e-
Acetophenone	<,J	8.0e0	<,J	1.1e1	<,J	1.1e1	<,J	2.3e2	<,J	3.8e-6	<,J	3.0e-
Aniline	<	2.4e1	<	3.4e1	<	3.4e1	<	6.7e2	<	1.1e-5	<	8.9e-
Anthracene	<	2.1e0	<	2.9e0	<	2.9e0	<	5.8e1	<	9.6e-7	<	7.6e-
Benzidine	<	1.4e2	<	2.0e2	<	2.1e2	<	4.1e3	<	6.8e-5	<	5.4e-
Benzoic acid	E		lε	8.2e2	Ε	8.3e2	E	1.6e4	Ε	2.7e-4	Ε	2.2e-
Benzo(a)anthracene	<	2.6e0	<	3.7e0	<	3.7e0	<	7.3e1	<	1.2e-6	<	9.7e-
Benzo(a)pyrene	<	2.8e1	<	4.0e1	<	4.0e1	<	7.9e2	<	1.3e-5	<	1.0e-
Benzo(b)fluoranthene	<	6.5e1	<	9.1e1	<	9.2e1	<	1.8e3	<	3.0e-5	<	2.4e-
Benzo(g,h,i)perylene	<	3.7e1	<	5.2e1	<	5.2e1	<	1.0e3	<	1.7e-5	<	1.4e-
Benzo(k)fluoranthene	<	9.3e1	<	1.3e2	<	1.3e2	<	2.6e3	<	4.4e-5	<	3.5e-
Benzyl alcohol	<	1.2e2	<	1.7e2	<	1.8e2	<	3.5e3	<	5.8e-5	<	4.6e-
ois(2-Chloroethoxy)methane	<	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1	<	1.0e-6	<	8.0e-
ois(2-Chloroethyl)ether	<	2.4e0	<	3.4e0	<	3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-
ois(2-Ethylhexyl)phthalate	<,J	3.5e1	<,J	4.9e1	<.J	4.9e1	<,J	9.7e2	<.J	1.6e-5	<, j	1.3e-
4-Bromophenyl-phenylether	<	2.0e0	<	2.9e0	<	2.9e0	<	5.7e1	<	9.5e-7	<	7.6e-
Butylbenzylphthalate	<	2.8e0	<	4.0e0	<	4.0e0	<	7.9e1	<	1.3e-6	<	1.0e-
Carbazole	<	2.8e0	<	4.0e0	<	4.0e0	<	7.9e1	<	1.3e-6	<	1.0e-
	<	3.7e0	<	5.2e0	<	5.2e0	<	1.0e2		1.7e-6		1.4e
4-Chloro-3-methylphenol	<	1.9e1	<	2.7e1	<	2.8e1	<	5.5e2		9.1e-6		7.2e
4-Chloroaniline	<	2.0e0	<	2.8e0	<	2.9e0	<		<	9.4e-7	<	7.5e-
2-Chloronaphthalene	<	2.4e0	<	3.4e0	<	3.4e0	<	6.7e1		1.1e-6		8.9e
2-Chlorophenol	<	2.4e0	<	3.4e0	<	3.4e0	<	6.7e1	<	1.1e-6		8.9e
4-Chlorophenyl phenyl ether	<		<	3.7e0	<	3.7e0	<	7.3e1	1.	1.2e-6		9.7e
Chrysene	1	2.6e0	1	3.7e0	<,J	3.1e0	<.J	6.1e2	l	1.0e-5		8.0e-
Di-n-butylphthalate	< J	2.2e1	<,J	4.9e1	1 '	4.9e1	1 '	9.7e2		1.6e-5	1 '	1.3e-
Di-n-octylphthalate	<,J	3.5e1	<,J		<,J	5.2e1	<,J <	1.0e3		1.7e-5		1.4e
Dibenz(a,h)anthracene	<	3.7e1	<	5.2e1 3.4e0	<	3.4e0	<	6.7e1		1.1e-6		8.9e
Dibenzofuran	<	2.4e0	<		<	3.4e0 3.4e0	<	6.7e1		1.1e-6	ı	8.9e
1,2-Dichlorobenzene	<	2.4e0	<	3.4e0	<	3.4e0 3.7e0	<	7.3e1	<	1.2e-6		9.7e
1,3-Dichlorobenzene	< .	2.6e0	<	3.7e0					ı			1.7e
1,4-Dichlorobenzene	<,J	4.5e0	<,J	6.4e0	<,J	6.5e0	<,J	1.3e2	_	2.1e-6		8.9e
3,3'-Dichlorobenzidine	<	2.4e1	<	3.4e1	<	3.4e1	<	6.7e2	< <	1.1e-5		1.0e-
2,4-Dichlorophenol	<	2.8e0	<	4.0e0	<	4.0e0	<	7.9e1	ŀ	1.3e-6	1	
Diethylphthalate	<	3.2e0	<	4.6e0	<	4.6e0	<	9.1e1		1.5e-6	1	1.2e
Dimethyl phthalate	<	2.1e0		2.9e0	<	3.0e0	<	5.8e1		9.7e-7		7.7e
2,4-Dimethylphenol	<	1.3e1	<	1.8e1	<	1.8e1	<	3.6e2	ł .	6.0e-6	1	4.7e
4,6-Dinitro-2-methylphenol	<	2.8e1	<	4.0e1	<	4.0e1	<	7.9 <b>e</b> 2		1.3e-5	1	1.0e
2,4-Dinitrophenol	<	5.8e1	<	8.2e1	<	8.3e1	<	1.6e3		2.7e-5	1	2.2e
2,4-Dinitrotoluene	<	2.8e0	<	4.0e0	<	4.0e0	<	7.9e1		1.3e-6	_	1.0e
2,6-Dinitrotoluene	<	2.6e0	<	3.7e0	<	3.7e0	<	7.3e1	1	1.2e-6		9.7e
1,2-Diphenylhydrazine	<	2.1e0	<	3.0e0	<	3.0e0	<	6.0e1	1	9.9e-7		7.9e
Fluoranthene	<	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1	1	1.0e-6		8.0e
Fluorene	<	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1		1.0e-6		8.0e
Hexachlorocyclopentadiene	<	3.5e1	<	4.9e1	<	4.9e1	<	9.7e2		1.6e-5		1.3e
Hexachlorobenzene	<	2.4e0	<	3.4e0	<	3.4e0		6.7 <b>e</b> 1		1.1e-6		8.9e
Hexachlorobutadiene	<	3.2e0	<	4.6e0	<	4.6e0	<	9.1e1	<	1.5e-6		1.2e
Hexachloroethane	< <	3.5e0	<	4.9e0	<	4.9e0	<	9.7e1	<	1.6e-6		1.3e
Indeno(1,2,3-cd)pyrene	<	3.2e1	<	4.6e1	<	4.6e1	<	9.1 <b>e</b> 2		1.5e-5		1.2e
Isophorone	<	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1		1.0e-6		8.0e
2-Methylnaphthalene	<	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1	<	1.0e-6		8.0e
2-Methylphenol	<	1.0e1	<	1.4e1	<	1.4e1	<	2.9e2	<	4.8e-6	<	3.8e

Table B-4. SVOC-END-2.

Run Date:

Project: 01-1062-01-0866

Run Identification: Run Type: Lab Report Date: Lab Report Status: (preliminary or final)

6/19/2001 0010-END-2 Test 8/28/2001 Final

### **RESULTS**

final)		WASS SLOW DATES										
		CON		TRATION:		MASS FLOW RATES						
	Actual			tandard		Standard						
	(µg/acm	)	()	µg/scm)	(µ	g/dscm)	μ	rg/min	gr	ams/sec	l	lb/h
3-Methylphenol & 4-Methylphenol	<	7.4e0	<	1.0e1	<	1.0e1	<	2.1e2	<	3.4e-6	<	2.7e-5
N-Nitroso-di-n-propylamine		2.4e0		3.4e0		3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-6
N-Nitrosodimethylamine		2.4e0	<	3.4e0		3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-6
N-Nitrosodiphenylamine		3.2e0	<	4.6e0	ł	4.6e0	<	9.1e1	<	1.5e-6	<	1.2e-5
Naphthalene		2.4e0	<	3.4e0		3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-6
2-Nitroaniline	< :	2.4e0	<	3.4e0	<	3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-6
3-Nitroaniline	<	3.9e0	<	1.2e1	<	1.3e1	<	2.5e2	<	4.2e-6		3.3e-5
4-Nitroaniline	<	7.8e0	<	1.1e1	<	1.1e1	<	2.2e2	<	3.7e-6	<	2.9e-5
Nitrobenzene	<,J :	2.6e0	<,J	3.7e0	ر >	3.7e0	<,J	7.3e1	<,J	1.2e-6	<,J	9.7e-6
2-Nitrophenol	1	1,1e1	<,J	1.6e1	<,J	1.6e1	<,J	3.1e2	<,J	5.2e-6	<,J	4.1e-5
4-Nitrophenol		1.2e1	<,J	1.7e1	<, J	1.7e1	<,J	3.3e2	<,J	5.6e-6	<,J	4.4e-5
2,2'-Oxybis(1-chloropropane)		3.0e0	<	4.3e0	<	4.3e0	<	8.5e1	<	1.4e-6	<	1.1e-5
Pentachlorobenzene	< :	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1	<	1.0e-6	<	8.0e-6
Pentachloronitrobenzene	< :	2.4e0	<	3.4e0		3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-6
Pentachlorophenol	< 1	3.0e1	<	1.1e2	<	1.1e2	<	2.3e3	<	3.8e-5	<	3.0e-4
Phenanthrene	1	2.1e0	<	3.0e0	<	3.0e0	<	6.0e1	<	9.9e-7	<	7.9e-6
Phenol	<,J	3.4e0	<,J	1.2e1	<,J	1.2e1	<,J	2.4e2	<,J	4.0e-6	<,J	3.1e-5
Pyrene	< :	2.2e0	<	3.0e0	<	3.1e0	<	6.1e1	<	1.0e-6	<	8.0e-6
Pyridine	< :	3.7e0	<	5.2e0	<	5.2e0	<	1.0e2	<	1.7e-6	<	1.4e-5
1,2,4,5-Tetrachlorobenzene	< :	2.4e0	<	3.4e0	<	3.4e0	<	6.7e1	<	1.1e-6	<	8.9e-6
1.2.4-Trichlorobenzene	<	2.6e0	<	3.7e0	<	3.7e0	<	7.3e1	<	1.2e-6	<	9.7e-6
2,4,5-Trichlorophenol	< !	5.4e0	<	7.6e0	<	7.7e0	<	1.5e2	<	2.5e-6	<	2.0e-5
2,4,6-Trichlorophenol	< :	3.5e0	<	4.9e0	<	4.9e0	<	9.7e1	<	1.6e-6	<	1.3e-5
TICs												
3-Hexanone	N,J,	1.5e1	N,J	2.1e1	N,J	2.2e1	N,J	4.3e2	N,J	7.1e-6	N,J	5.6e-5
Benzaldehyde		1.6e2	N,J	2.2e2	N,J	2.2e2	N,J	4.4e3	N,J	7.4e-5	N,J	5.9e-4
2-Cyclohexene-1-one, 3-methyl-		.9e-1	N,J	9.7e-1	N,J	9.8e-1	N,J	1.9e1	N,J	3.2e-7	N,J	2.6e-6
Formic acid, phenylmethyl ester		2.1e1		2.9e1	N,J	2.9e1	N,J	5.8e2	N,J	9.6e-6	N,J	7.6e-5
Benzaldehyde, ethyl-	N,J,	1.3e1	N,J	1.9e1	N,J	1.9e1	N,J	3.7e2	N,J	6.2e-6	N,J	4.9e-5
Dodecane		3.0e0	N,J	1.1e1	N,J	1.1e1	N,J	2.3e2	N,J	3.8e-6	N,J	3.0e-5
Tridecane		4.5e0	N,J	6.4e0	N,J	6.5e0	N,J	1.3e2	N,J	2.1e-6	N,J	1.7e-5
2,4-Hexadiene	N,J, €	3.1e0	N,J	8.5e0	N,J	8.6e0	N,J	1.7e2	N,J	2.8e-6	N,J	2.3e-5
2,5-Diethylphenol	N,J, 2	2.2e1	N,J	3.0e1	N,J	3.1e1	N,J	6.1e2	N,J	1.0e-5	N,J	8.0e-5
Tetradecane	N,J, 2	2.1e1	N,J	3.0e1	N,J	3.0e1	N,J	6.0e2	N,J	1.0e-5	N,J	8.0e-5
Hexatriacontane	N,J,	1.2e0	N,J	1.7e0	N,J	1.8e0	N,J	3.5e1	N,J	5.8e-7	N,J	4.6e-6
Phosphoric acid tributyl ester	N,J, :	5.2e0	N,J	7.3e0	N,J	7.4e0	N,J	1.5e2	N,J	2.4e-6	N,J	1.9e-5
Cyclododecane	N,J, 2	2.8e0	N,J	4.0e0	N,J	4.0e0	N,J	7.9e1	N,J	1.3e-6	N,J	1.0e-5
Pentadecane	N,J, 9	.7e-1	N,J	1.4e0	N,J	1.4e0	N,J	2.7e1	N,J	4.6e-7	N,J	3.6e-6
Heneicosane	N,J, :	2.0e0	N,J	2.8e0	N,J	2.8e0	N,J	5.5e1	N,J	9.2e-7	N,J	7.3e-6
Tetracosane	N,J, -	5.6e0	N,J	7.9e0	N,J	8.0e0	N,J	1.6e2	N,J	2.6e-6	N,J	2.1e-5
Pentacosane	N,J, 8	3.0e0	N,J	1.1e1	Ñ,Ĵ	1.1e1	N,J	2.3e2	N,J	3.8e-6	N,J	3.0e-5
Hexacosane	N,J,	1.4e1	N,J	1.9e1	N,J	2.0e1	N,J	3.9e2	N,J	6.5e-6	N,J	5.2e-5
Heptacosane	N,J,	1.8e1	N,J	2.5e1	N,J	2.6e1	N,J	5.1e2	N,J	8.4e-6	N,J	6.7e-5
Hexatriacontane	N,J, 4	1.3e1	N,J	6.1e1	N,J	6.2e1	N,J	1.2e3	N,J		N,J	1.6e-4
Eicosane	N,J, 4	1.1e0	N,J	5.8e0	N,J	5.8e0	N,J	1.2e2	N,J	1.9e-6	N,J	1.5e-5
Furan, 2,5-dimethyl-												
Heptane, 2,5-dimethyl-				į								
Heptane, 2,3-dimethyl-												
Benzoic acid, methyl ester					"							
Heptadecane												
Octodecane												
Phosphine oxide, triphenyl-												
Nonacosane												
Tetratriacontane					L			,,				
- Ciratriacontante	<u> </u>											

Table B-5. 0031-STRT-1.							
VOST SAMI	PLING DAT	A SHEF	<u></u>				
Site: HLLWE Offgas Tie-in Run No.:	0031-S1		Meter Box N	lo.:	1 & 2		
Project: 01-1062-01-0866 Run Type:	Tes		Y-factor:			4/1.005	
Date: 6/20/2001 Pbar., in. Hg:	25.29		Operator:			/FE/JA	
Date. 0/20/2001   Foar., III. Fig.	20.2	<u> </u>	Toberaior.		1 \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	,	
VOST Tube Leak Check Sampling Sar	mpling	Probe	Condenser	Meter	Meter	Pump	
	ime	Temp.	Temp.	Temp.	Volume	Vacuum	
Numbers (L/min) (in. Hg) (L/min) (24 hr)	(min.)	(°C)	(°C)	(°C)	(L)	(in. Hg)	
		130	5.0	15	0.000	4.5	
		130	5.0	17	2.884	4.5	
I TOTAL CONTRACTOR OF THE CONT	description of the second	130	4.0	17	6.083	4.5	
		130	5.0	19	9.194	4.7	
A-3361 8:25 8:30		130	5.0	19	12.439	4.7	
8:35	25	130	5.0	21	15.653	4.7	
8:35	30	130	5.0	21	18.725	4.7	
		130	6.0	22	21.483	4.3	
8:45 8:50	decision of the second	130	6.0	22	24.230	4.3	
Post-test 8:50		130	0.0		24.230	Max:	
0.003 5.5 Total 0:40	40	130	5.1	19	∠4.∠3U	Max: 4.7	
Average					0.000	7.5	
Set 2 Pretest Target: 9:00		130	7.0	22	0.000	7.5 11.0	
A-3362 0.007 16 1L/MIN 9:05		130	6.0	23	3.574		
A-3363 9:10		130	6.0	24	8.376	12.5 12.5	
A-3364 9:15		130	6.0	26	13.373		
9:20		130	8.0	25	18.381	12.5	
9:25	25	130	8.0	25	23.393	12.5	
Post-test 9:26		130	8.0	25	24.020	12.5	
0.007 16 Total 0:26	26	400	<b> </b>	0.4	24.020	Max:	
Average	-	130	7.0	24		12.5	
Set 3 Pretest Target: 9:48		130	10.0	27	0.000	9.5	
A-3365 0.007 16 9:53		130	8.0	28	5.031	9.5	
A-3366 9:58		130	8.0	28	10.566	10.0	
A-3337 10:03		130	8.0	28	15.901	9.0	
10:08		130	8.0	29	21.074	9.0	
Post-test 10:11		130	8.0	29	24.202	9.0	
0.004 16 Total 0:23	23	400			24.202	Max:	
Average		130	8.3	28		10.0	
Set 4         Pretest         Target:         10:23           10:23         10:28         10:28		130	13.0	29	0.000	12.7	
A-3368 0.005 16 10:28		130	9.0	29	4.833	12.7	
A-3369 10:33		130	7.0	30	9.742	12.7	
A-3370 10:38		130	7.0	30	14.662	12.7	
10:43		130	7.0	30	19.598	12.8	
Post-test 10:48		130	7.0	31	24.544 24.544	12.8	
0.002 16 Total 0:25	25				. /4.5441	Max:	
	(2003)	400	0.0	20		40.0	
Average	L	130	8.3	30	2.1011	12.8	
Condensate Average Tenax R	L	Anasor	8.3 b Rinse:	30	21.011	12.8	
Condensate A-3371 Average Tenax R	linse:	Anasorl NA	b Rinse:	30		12.8	
Condensate A-3371 NA Final Condensate Volume: Average	Rinse: inse Vol.:	Anasorl NA Anasorb	b Rinse:  Rinse Vol.:	30		12.8	
Condensate Tenax R A-3371 NA Final Condensate Volume: Tenax R 40 mL NA	Rinse: linse Vol.; mL	Anasorl NA Anasorb NA	b Rinse:	30		12.8	
Condensate A-3371 Final Condensate Volume: 40 mL Average  Tenax R NA Tenax R NA ADDIT	Rinse: inse Vol.: mL TONAL INP	Anasorl NA Anasorb NA UTS	b Rinse: Rinse Vol.: mL			America View Line - Williams	
Condensate A-3371 Final Condensate Volume: 40 mL ADDIT Symbol	kinse: inse Vol.: mL IONAL INP Units	Anasorl NA Anasorb NA UTS Set 1	b Rinse:  Rinse Vol.:  mL  Set 2	Set 3	Set 4	Average	
Average   Average	Rinse: inse Vol.: mL TONAL INP Units %	Anasorl NA Anasorb NA UTS Set 1 20.5	Rinse Vol.: mL Set 2 20.5	Set 3 20.5	Set 4 20.5	Average 20.5	
Average   Average	inse Vol.: mL TONAL INP Units % dscm/sec	Anasorb NA Anasorb NA UTS Set 1 20.5 0.330	b Rinse:  Rinse Vol.: mL  Set 2  20.5 0.330	Set 3	Set 4 20.5	Average 20.5	
Average   Average	inse Vol.: mL TONAL INP Units % dscm/sec	Anasori NA Anasorb NA UTS Set 1 20.5 0.330 PARAMI	Rinse Vol.: mL Set 2 20.5 0.330 ETERS	Set 3 20.5 0.330	Set 4 20.5 0.330	Average 20.5 0.330	
Average   Average	inse Vol.:  mL  TONAL INP  Units  dscm/sec  AMPLING I	Anasori NA Anasori NA UTS Set 1 20.5 0.330 PARAMI Set 1	Set 2 20.5 0.330 ETERS Set 2	Set 3 20.5 0.330 Set 3	Set 4 20.5 0.330	Average 20.5 0.330 Net	
Average   Average	inse Vol.: mL TONAL INP Units % dscm/sec	Anasori NA Anasorb NA UTS Set 1 20.5 0.330 PARAMI	Set 2 20.5 0.330 ETERS Set 2	Set 3 20.5 0.330	Set 4 20.5 0.330	Average 20.5 0.330	
Average   Average	inse Vol.: mL IONAL INP Units % dscm/sec AMPLING I Units dsL	Anasori NA Anasori NA UTS Set 1 20.5 0.330 PARAME Set 1 20.508	Set 2 20.5 0.330 ETERS Set 2 20.116	Set 3 20.5 0.330 Set 3 20.008	Set 4 20.5 0.330 Set 4 20.179	Average 20.5 0.330  Net 80.810	
Average   Average	inse Vol.:  mL  TONAL INP  Units  dscm/sec  AMPLING I	Anasori NA Anasori NA UTS Set 1 20.5 0.330 PARAMI Set 1	Set 2 20.5 0.330 ETERS Set 2 20.116	Set 3 20.5 0.330 Set 3 20.008	Set 4 20.5 0.330	Average 20.5 0.330 Net	

Table B-5. 0031-STRT-1.			T			
1401C B-3. 0031-311(1-1.	CC	NCENTRATIONS				
		er dry standard cub	oic meter			
Project:	0866	Lab Report Date:	08/20/01			
Run Date:	6/20/2001	Lab Report Status:	Final			
Run Identification:	0031-STRT-1					
Analyte						n Total
	<del></del>				Flag B	μg/dscm 8.9e1
Acetone					<u> </u>	3.5e1
Acrylonitrile						
Benzene			MARKY		<	4.9e0
Bromobenzene	5.5.10.5.5.40.000 (ATT)					1.5e0
Bromochloromethane		ne at 1 to P / P				1.9e0
Bromodichloromethane					<	1.5e0
Bromoform					<u> </u>	2.2e0
Bromomethane					<,J	2.1e0
2-Butanone					<,J	1.1e1
n-Butylbenzene						1.9e0
sec-Butylbenzene					<	1.1e0
tert-Butylbenzene					<	1.7e0
Carbon disulfide					<	1.2e1
Carbon tetrachloride					<	2.0e0
Chlorobenzene					<	1.2e0
Chlorodibromomethane					<	1.9e0
Chloroethane	THE STATE OF THE S				<,J	2.0e0
Chloroform					<,J	3.0e0
Chloromethane					J	1.5e1
2-Chlorotoluene					<	7.3e-1
4-Chlorotoluene						7.3e-1
1,2-Dibromo-3-chloropropane					<	3.3e0
1,2-Dibromoethane	,				<	2.5e0
Dibromomethane					<	2.1e0
1,2-Dichlorobenzene					<	2.2e0
1,3-Dichlorobenzene					<	1.2e0
1,4-Dichlorobenzene					<	1.7e0
Dichlorodifluoromethane		- 245 A - 5			_ <	3.2e0
1,1-Dichloroethane					<	1.9e0
1,2-Dichloroethane				,,	<,J	2.0e0
1,1-Dichloroethene					<,J	2.0e0
cis-1,2-Dichloroethene					<	1.9e0
trans-1,2-Dichloroethene		. 41 - 1000 1100 1100 1100 1100 1100 1100			<	2.1e0
1,2-Dichloropropane					<	1.6e0
1,3-Dichloropropane					<	2.1e0
2,2-Dichloropropane					<	2.0e0
1,1-Dichloropropene					<	2.2e0
cis-1,3-Dichloropropene					<	1.6e0
trans-1,3-Dichloropropene					<	1.9e0
Ethylbenzene					<	1.2e0
Hexachlorobutadiene				,	<	2.7e0
2-Hexanone					<	6.9e0
Isopropylbenzene					<	8.7e-1
p-Isopropyltoluene					<	1.4e0
Methylene chloride					E,B	2.2e2
4-Methyl-2-pentanone			······································	-	<	7.3e0
Naphthalene					<	2.6e0
n-Propylbenzene					<	8.3e-1

Table B-5. 0031-STRT-1.					
		NCENTRATION per dry standar	meter	 -	
Project:	0866	Lab Report Dat	08/20/01		
Run Date:	6/20/2001	Lab Report Sta	Final		
Run Identification:	0031-STRT-1	,			
Analyte				 l l	n Total
			 	 Flag	μg/dscm
Styrene			 	 <	9.7e-1
1,1,1,2-Tetrachloroethane			 	 <	1.2e0
1,1,2,2-Tetrachloroethane			 	 <	2.7e0
Tetrachloroethene			 	 <	1.9e0
Toluene			 	 <,J	2.4e0
1,2,3-Trichlorobenzene			 	 <	2.6e0
1,2,4-Trichlorobenzene			 	 <	2.7e0
1,1,1-Trichloroethane			 	 <	2.4e0
1,1,2-Trichloroethane			 	 <	2.1e0
Trichloroethene			 	 <	2.0e0
Trichlorofluoromethane			 	 <,J	2.0e0
1,2,3-Trichloropropane			 ***	 <	3.0e0
1,2,4-Trimethylbenzene			 	 <	1.2e0
1,3,5-Trimethylbenzene			 	<	7.3e-1
Vinyl chloride			 	 <,J	1.6e0
m-Xylene & p-Xylene			 	 <	5.4e0
o-Xylene			 	 <	9.5e-1
TICS			 	 	
Hexane, 2-methyl-			 	 N,J,M	2.1e0
Pentane, 2,3-dimethyl-				 N,J,M	2.2e0
Butane, 1-chloro-	***		 	 N,J,M	
Hexane, 3-methyl-	10 10 10 10 10 10 10 10 10 10 10 10 10 1		 	 N,J,M	4.7e0
Cyclohexene			 	 N,J,M	1.2e0
1-Heptene				N,J,M	6.7e-1
Cyclohexane, methyl-			 	 N,J,M	1.4e0
Hexane, 2,4-dimethyl-			 	 N,J,M	1.4e0
Cyclopentane, ethyl-			 	 N,J,M	3.5e-1
Octane				 N,J,M	3.3e-1
Decane			 	 N,J,M	1.5e0
Undecane			 	 N,J,M	1.2e1
Undecane, 5-methyl-		AA - METTER	 	 N,J,M	7.9e0
Decane, 2,9-dimethyl-				 N,J,M	7.9e-1
Dodecane			 	 N,J,M	3.8e2
Undecane, 2,6-dimethyl-			 	 N,J,M	1.4e0
Cyclohexane, hexyl-				N,J,M	7.3e-1
Tridecane				N,J,M	3.5e1
Tetradecane				 N,J,M	1.1e1
Pentane, 3,3-dimethyl-			 		
Pentane, 3-ethyl-			 	 	
Cyclopentane, 1,2-dimethyl-					

Table B-11. 0031-STRT-1	MASS FLOW RATE		
	grams per second		
Project:	01-1062-01-086 Lab Report Date: 08/20/01		
Run Date:	6/20/2001 Lab Report Status: Final		
Run Identification:	0031-STRT-1		<del></del>
Analyte		Flag	n Total g/sec
A 1		B	2.9e-5
Acetone		<u> </u>	1.1e-5
Acrylonitrile			1.6e-6
Benzene			4.9e-7
Bromobenzene			6.1e-7
Bromochloromethane			4.9e-7
Bromodichloromethane			7.3e-7
Bromoform			6.9e-7
Bromomethane		<,J	3.6e-6
2-Butanone		<u>``,</u> `	6.1e-7
n-Butylbenzene			3.5e-7
sec-Butylbenzene		<	5.7e-7
tert-Butylbenzene			
Carbon disulfide			4.0e-6
Carbon tetrachloride		<	6.5e-7
Chlorobenzene		<u> </u>	3.9e-7
Chlorodibromomethane		<u> </u>	6.1e-7
Chloroethane		<,J	6.5e-7
Chloroform		<,J	9.8e-7
Chloromethane		J	4.9e-6
2-Chlorotoluene			2.4e-7
4-Chlorotoluene		. <	2.4e-7
1,2-Dibromo-3-chloropropane	)	<	1.1e-6
1,2-Dibromoethane		<	8.2e-7
Dibromomethane		<	6.9e-7
1,2-Dichlorobenzene		<	7.3e-7
1,3-Dichlorobenzene		<	4.1e-7
1,4-Dichlorobenzene		<	5.7e-7
Dichlorodifluoromethane		<	1.1e-6
1,1-Dichloroethane		<	6.1e-7
1,2-Dichloroethane		<,J	6.5e-7
1,1-Dichloroethene		<,J	6.5e-7
cis-1,2-Dichloroethene		<	6.1e-7
trans-1,2-Dichloroethene		<	6.9e-7
1,2-Dichloropropane		<	5.3e-7
1,3-Dichloropropane		<	6.9e-7
2,2-Dichloropropane		<	6.5e-7
1,1-Dichloropropene		<	7.3e-7
cis-1,3-Dichloropropene		<	5.3e-7
trans-1,3-Dichloropropene		<	6.1e-7
Ethylbenzene		<	3.8e-7
Hexachlorobutadiene		<	9.0e-7
2-Hexanone		<	2.3e-6
Isopropylbenzene		<	2.9e-7
p-Isopropyltoluene		<	4.5e-7
Methylene chloride		E,B	7.3e-5
4-Methyl-2-pentanone		<	2.4e-6
Naphthalene		<	8.6e-7
n-Propylbenzene		<	2.7e-7

Table B-11. 0031-STRT-1	.   !				
THE DITTO OF STREET		ASS FLOW RATE	L:		
		rams per second			
Project:		Lab Report Date:	08/20/01		
Run Date:	6/20/2001	Lab Report Status:	Final		
Run Identification: Analyte	0031-STRT-1			Rur	Total
Analyte				Flag	g/sec
Styrene				<	3.2e-7
1,1,1,2-Tetrachloroethane				<	4.0e-7
1,1,2,2-Tetrachloroethane	• • • • • • • • • • • • • • • • • • • •			<	9.0e-7
Tetrachloroethene				<	6.1e-7
Toluene				<,J	7.8e-7
1,2,3-Trichlorobenzene				<	8.6e-7
1,2,4-Trichlorobenzene				<	9.0e-7
1,1,1-Trichloroethane				<	7.8e-7
1,1,2-Trichloroethane				<	6.9e-7
Trichloroethene				<	6.5e-7
Trichlorofluoromethane				<,J	6.5e-7
1,2,3-Trichloropropane				<	9.8e-7
1,2,4-Trimethylbenzene				<	4.1e-7
1,3,5-Trimethylbenzene				<	2.4e-7
Vinyl chloride				<,J	5.3e-7
m-Xylene & p-Xylene				<	1.8e-6
o-Xylene				<	3.1e-7
TICS					
Hexane, 2-methyl-	All Ale The State of the State			N,J,M	6.9e-7
Pentane, 2,3-dimethyl-		, ,, , ,		N,J,M	7.3e-7
Butane, 1-chloro-				N,J,M	2.3e-7
Hexane, 3-methyl-				N,J,M	1.6e-6
Cyclohexene				N,J,M	4.1e-7
1-Heptene				N,J,M	2.2e-7
Cyclohexane, methyl-				N,J,M	4.5e-7
Hexane, 2,4-dimethyl-				N,J,M	4.5e-7
Cyclopentane, ethyl-				N,J,M	1.1e-7
Octane				N,J,M	1.1e-7
Decane				N,J,M	4.9e-7
Undecane				N,J,M	4.1e-6
Undecane, 5-methyl-				N,J,M	2.6e-6
Decane, 2,9-dimethyl-				N,J,M	2.6e-7
Dodecane				N,J,M	1.3e-4
Undecane, 2,6-dimethyl-				N,J,M	4.5e-7
Cyclohexane, hexyl-				N,J,M	2.4e-7
Tridecane				N,J,M	1.1e-5
Tetradecane				N,J,M	3.8e-6
Pentane, 3,3-dimethyl-					
Pentane, 3-ethyl-					
Cyclopentane, 1,2-dimethyl-					

Site: Project:		ID-1.	İ		i i					
Project:			VO	ST SAM	LING DAT	A SHEE	Т			•
Project:	HIIWE (	Offgas Tie-in			0031-E		Meter Box N	lo.:		1
	01-1062-		Run Type:		Te		Y-factor:	-	0.	998
Date:		/20/01	Pbar., in. H	a.	25.2		Operator:			FE,JA
Date.		20701	11 0011, 1111	9'			1-1-1-1-1			
VOST Tube	Leak	Check	Sampling	Sar	npling	Probe	Condenser	Meter	Meter	Pump
Sample	Rate	@ vacuum	Rate		ime	Temp.	Temp.	Temp.	Volume	Vacuum
Numbers	(L/min)	(in. Hg)	(L/min)	(24 hr)	(min.)	(°C)	(°C)	(°C)	(L)	(in. Hg)
Set 1		etest	Target:	14:00	0	130	11.0	31	0.000	13.0
A-3409	0.001	16	1		5	130	10.0	31	4.788	13.0
A-3410	0.001			14:10	10	130	10.0	31	9.586	13.0
A-3411				14:15	15	130	11.0	33	14.413	13.0
A-3411				14:20	20	130	11.0	33	19.260	13.0
		st-test		14:25	25	130	11.0	34	24.164	13.0
	0.003		Total	0:25		100		<u> </u>	24.164	Max:
	0.003	10	Average	0.20		130	10.7	32		13.0
C-4.7	D.	etest	Target:	14:36	0	130	10.0	34	0.000	12.0
Set 2	0.000	16	1 1			130	10.0	35	4.994	12.0
A-3412	0.000_	10	ļ <u>'</u>	14:41	10	130	10.0	35	10.001	12.0
A-3413				14:46	15	130	10.0	35	15.128	12.0
A-3414		! <del> </del>	j —	14:51	20	130	10.0	36	20.296	12.0
l		st-test		15:00	24	130	10.0	35	24.452	12.0
			T-4-1			130	10.0	33	24.452	Max:
	0.008	16	Total	0:24		130	10.0	35	24.402	12.0
<u> </u>		11	Average	45.40	^	130	12.0	35 35	0.000	5.0
Set 3		etest	Target:	15:10	0					5.0
A-3415	0.001	16	1		5	130	11.0	35	1.909	
A-3416				15:20	10	130	11.0	35	3.721	5.0
A-3417				15:25	15	130	11.0	35	5.650	5.0
				15:30	20	130	11.0	35	7.605	5.0
				15:35	25	130	11.0	35	9.514	5.0
		· +		15:40	30	130	11.0	35	11.410	5.0
				15:45	35	130	11.0	35	13.348	5.0
				15:50	40	130	11.0	35	15.321	5.0
				15:55	45	130	11.0	35	17.238	5.0
				16:00	50	130	11.0	35	19.154	5.0
		<u> </u>		16:05	55	130	11.0	35	21.118	5.0
		st-test		16:10	60	130	12.0	35	23.165	5.0
	0.000	6		16:15	65	130	12.0	35	24.206	5.0
			Total	1:05	65				24.206	Max:
			Average			130	11.2	35		5.0
Set 4	l .	etest	Target:	16:29	0	130	11.0	31	0.000	4.0
A-3418	0.000	5	1		5	130	11.0	31	2.646	4.0
A-3419				16:39	10	130	11.0	31	5.400	4.0
A-3420		ı	1	16:44	1	130	11.0	32	7.965	4.0
				16:49		130	11.0	33	10.607	4.0
				16:54		130	11.0	33	13.270	4.0
1		ļ <u></u>	<u> </u>	16:59		130	11.0	34	15.873	4.0
1			1	17:04	35	130	11.0	34	18.500	4.0
				17:09	40	130	12.0	34	21.135	4.0
		st-test		17:09 17:14	40 45	130 130	12.0 11.0	34 35	21.135 23.825	4.0 4.0
	Po: 0.000	st-test 5		17:09 17:14 17:15	40 45 46	130	12.0	34	21.135 23.825 24.404	4.0 4.0 4.0
			Total	17:09 17:14	40 45 46	130 130	12.0 11.0 11.0	34 35 35	21.135 23.825	4.0 4.0 4.0 Max:
			Total Average	17:09 17:14 17:15 0:46	40 45 46 46	130 130 130 130	12.0 11.0 11.0	34 35	21.135 23.825 24.404	4.0 4.0 4.0
Condensate	0.000			17:09 17:14 17:15	40 45 46 46	130 130 130 130	12.0 11.0 11.0	34 35 35	21.135 23.825 24.404	4.0 4.0 4.0 Max:
Condensate A-3421	0.000			17:09 17:14 17:15 0:46	40 45 46 46	130 130 130 130 Anasori NA	12.0 11.0 11.0 11.1 5 Rinse:	34 35 35	21.135 23.825 24.404	4.0 4.0 4.0 Max:
	0.000	5		17:09 17:14 17:15 0:46 Tenax R NA Tenax R	40 45 46 46 <b>Rinse:</b>	130 130 130 130 Anasori NA Anasorb	12.0 11.0 11.0 11.1 b Rinse:	34 35 35	21.135 23.825 24.404	4.0 4.0 4.0 Max:
A-3421 Final Conder	0.000	5		17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA	40 45 46 46 Rinse:	130 130 130 130 Anasori NA Anasorb NA	12.0 11.0 11.0 11.1 5 Rinse:	34 35 35	21.135 23.825 24.404	4.0 4.0 4.0 Max:
A-3421 Final Conder	0.000	5		17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA	40 45 46 46 <b>Rinse:</b>	130 130 130 130 Anasori NA Anasorb NA	12.0 11.0 11.0 11.1 b Rinse:	34 35 35 33	21.135 23.825 24.404 24.404	4.0 4.0 4.0 Max: 4.0
A-3421 Final Conder	0.000	5		17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA	40 45 46 46 Linse: inse Vol.: mL IONAL INP	130 130 130 130 Anasorl NA Anasorb NA UTS Set 1	12.0 11.0 11.0 11.1 5 Rinse: 9 Rinse Vol.: mL	34 35 35 33 33	21.135 23.825 24.404 24.404	4.0 4.0 4.0 Max:
A-3421 Final Conder	0.000	5 ume:	Average	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol Co2	40 45 46 46 kinse: inse Vol.: mL IONAL INP Units %	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5	12.0 11.0 11.0 11.1 5 Rinse: 9 Rinse Vol.: mL Set 2 20.5	34 35 35 33 33 Set 3 20.5	21.135 23.825 24.404 24.404 Set 4 20.5	4.0 4.0 4.0 Max: 4.0 Average 20.5
A-3421 Final Conder 40	0.000  nsate VolumL  Oxygen 0	5 ume:	Average	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol	40 45 46 46 Linse: inse Vol.: mL IONAL INP	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5	12.0 11.0 11.0 11.1 5 Rinse: 9 Rinse Vol.: mL Set 2 20.5	34 35 35 33 33	21.135 23.825 24.404 24.404 Set 4 20.5	4.0 4.0 4.0 Max: 4.0
A-3421 Final Conder 40 Average Dry	0.000  nsate VolumL  Oxygen 0	5 ume:	Average	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol Co2 Qsd	40 45 46 46 kinse: inse Vol.: mL IONAL INP Units %	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5 0.330	12.0 11.0 11.0 11.1 b Rinse: c Rinse Vol.: mL Set 2 20.5 0.330	34 35 35 33 33 Set 3 20.5	21.135 23.825 24.404 24.404 Set 4 20.5	4.0 4.0 4.0 Max: 4.0 Average 20.5
A-3421 Final Conder 40 Average Dry	0.000  nsate VolumL  Oxygen 0	5 ume:	Average	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol Co2 Qsd	40 45 46 46  Linse: inse Vol.: mL IONAL INP Units % dscm/sec	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5 0.330	12.0 11.0 11.0 11.1 b Rinse: c Rinse Vol.: mL Set 2 20.5 0.330	34 35 35 33 33 Set 3 20.5	21.135 23.825 24.404 24.404 Set 4 20.5 0.330	4.0 4.0 4.0 Max: 4.0 Average 20.5
A-3421 Final Conder 40 Average Dry Process Gas F	0.000  nsate VolumL  Oxygen ( Flow (dry, §	5 ume: Concentratio	Average	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol Co2 Qsd	40 45 46 46  Linse: inse Vol.: mL IONAL INP Units % dscm/sec	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5 0.330 PARAME	12.0 11.0 11.0 11.1 b Rinse: D Rinse Vol.: mL Set 2 20.5 0.330 ETERS	34 35 35 33 33 Set 3 20.5 0.330	21.135 23.825 24.404 24.404 Set 4 20.5 0.330	4.0 4.0 4.0 Max: 4.0 Average 20.5 0.330
A-3421 Final Conder 40 Average Dry	nsate VolumL Oxygen (Grow (dry, State and Stat	5  Concentration STP) @ 68°F  andard Concentration	Average	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol Co2 Qsd _ATED S Symbol	40 45 46 46  linse:  inse Vol.: mL  IONAL INP  Units % dscm/sec  AMPLING  Units	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5 0.330 PARAME	12.0 11.0 11.0 11.1 b Rinse: D Rinse Vol.: mL Set 2 20.5 0.330 ETERS	34 35 35 33 33 Set 3 20.5 0.330	21.135 23.825 24.404 24.404 Set 4 20.5 0.330	4.0 4.0 4.0 Max: 4.0 Average 20.5 0.330
A-3421 Final Conder 40  Average Dry Process Gas F	0.000  nsate VolumL  Oxygen ( Flow (dry, \$  ume @ Starry*Pbar*V	5  Concentration STP) @ 68°F  andard Concentration Concent	Average  CALCUI	17:09 17:14 17:15 0:46 Tenax R NA Tenax R NA ADDIT Symbol Co2 Qsd _ATED S Symbol	40 45 46 46  linse:  inse Vol.: mL  IONAL INP  Units % dscm/sec  AMPLING  Units	130 130 130 130 Anasort NA Anasort NA UTS Set 1 20.5 0.330 PARAME	12.0 11.0 11.0 11.1 b Rinse: D Rinse Vol.: mL Set 2 20.5 0.330 ETERS Set 2 19.629	34 35 35 33 33 Set 3 20.5 0.330	21.135 23.825 24.404 24.404 Set 4 20.5 0.330	4.0 4.0 4.0 Max: 4.0 Average 20.5 0.330 Net 78.355

Table B-6. 0031-END-1.		NOENTRATIONS			
		NCENTRATIONS er dry standard cubi	ic meter		
Project:	01-1062-01-	Lab Report Date:	08/20/01		
Run Date:		Lab Report Status:	Final		
Run Identification:	0031-END-1			Г Б	n Total
Analyte				Flag	n rotai μg/dscn
Acetone		**************************************		В	9.2e1
Acrylonitrile					3.6e1
Benzene				- ·	3.1e0
Bromobenzene				<	1.5e0
Bromochloromethane				<	1.9e0
Bromodichloromethane				<	1.5e0
Bromoform				<	2.3e0
Bromomethane				<,J	2.9e0
2-Butanone				,,, <,J	1.1e1
n-Butylbenzene			own e	<	1.9e0
sec-Butylbenzene		··· ·· · · - · · · · · · · · · · · · ·		<	1.1e0
tert-Butylbenzene	***************************************			<	1.8e0
Carbon disulfide	Committee and the state of the			<	1.5e1
Carbon tetrachloride				<	2.0e0
Chlorobenzene	777 - 1870/1988			<,J	1.2e0
Chlorodibromomethane				<	1.9e0
Chloroethane			- Address and Address Conf. (Market Conf.) (1) (1) (1) (1) (1) (1) (1) (1)	<,J	2.6e0
Chloroform				<	5.1e0
Chloromethane				<	3.7e1
2-Chlorotoluene				<	7.5e-1
4-Chlorotoluene				<	7.5e-1
1,2-Dibromo-3-chloropropane				<	3.6e0
1,2-Dibromoethane				<	2.6e0
Dibromomethane				<	2.2e0
1,2-Dichlorobenzene				<	2.3e0
1,3-Dichlorobenzene				<	1.3e0
1,4-Dichlorobenzene				<	1.8e0
Dichlorodifluoromethane				<,J	2.0e0
1,1-Dichloroethane				<	1.9e0
1,2-Dichloroethane				<,J	2.0e0
1,1-Dichloroethene				<,J	2.3e0
cis-1,2-Dichloroethene				<	1.9e0
trans-1,2-Dichloroethene				<	2.0e0
1,2-Dichloropropane		Mar. 1		<,J	1.7e0
1,3-Dichloropropane				<	2.2e0
2,2-Dichloropropane				<	2.0e0
1,1-Dichloropropene				<	2.3e0
cis-1,3-Dichloropropene				<	1.7e0
trans-1,3-Dichloropropene				<	1.9e0
Ethylbenzene				<	1.2e0
Hexachlorobutadiene				<	2.8e0
2-Hexanone				<	7.1e0
Isopropylbenzene				<	8.9e-1
p-Isopropyltoluene				<	1.4e0
Methylene chloride				В	2.2e1
4-Methyl-2-pentanone				<<	7.5e0
Naphthalene				<	2.7e0

Table B-6. 0031-END-1.					
· ·		ONCENTRATIONS		'	1
		per dry standard cub			
Project:	01-1062-01-	Lab Report Date:	08/20/01		
Run Date: Run Identification:	6/20/2001 0031-END-1	Lab Report Status:	Final		
Analyte	003 I-END-1	<u></u>		Rui	n Total
Analyte				Flag	μg/dscm
n-Propylbenzene				<	8.4e-1
Styrene				<	1.0e0
1,1,1,2-Tetrachloroethane				<	1.3e0
1,1,2,2-Tetrachloroethane				<	2.8e0
Tetrachloroethene				<	1.9e0
Toluene				<,J	3.6e0
1,2,3-Trichlorobenzene		mana Man of Miles of a Communication of the Communi		<	2.7e0
1,2,4-Trichlorobenzene				<	2.8e0
1,1,1-Trichloroethane				<	2.4e0
1,1,2-Trichloroethane				<	2.2e0
Trichloroethene			A THE PARTY OF THE	<	2.0e0
Trichlorofluoromethane				<,J	2.0e0
1,2,3-Trichloropropane				<	3.1e0
1,2,4-Trimethylbenzene				<	1.3e0
1,3,5-Trimethylbenzene				<	7.5e-1
Vinyl chloride				<,J	2.3e0
m-Xylene & p-Xylene				<	5.6e0
o-Xylene				<,J	1.0e0
TICs					
Pentane, 3,3-dimethyl-				N,J,M	7.5e-1
Hexane, 2-methyl-				N,J,M	4.0e0
Pentane, 2,3-dimethyl-		As Adribu V 1 18 Lores, your generality page species or species or species.		N,J,M	4.0e-1
Hexane, 3-methyl-				N,J,M	9.7e0
Cyclohexene				N,J,M	4.6e-1
Cyclobutane, ethenyl-				N,J,M	1.5e0
Cyclopentane, 1,2-dimethyl-, t		ALL SECTION AND THE SECTION AN		N,J,M	1.0e0
Cyclohexane, methyl-				N,J,M	2.6e0
Hexane, 2,4-dimethyl-				N,J,M	2.3e0
Cyclopentane, ethyl-				N,J,M	5.2e-1
Methane, trichloronitro-				N,J,M	4.6e0
Benzonitrile				N,J,M	2.4e0
Undecane				N,J,M	6.4e0
Undecane, 5-methyl-			<del></del>	N,J,M	2.3e0
Dodecane	Martin Address Charles Charles Co.	A BANK IN A BANK CHIEF CONTROL OF THE CONTROL OF TH		N,J,M	3.3e2
Tridecane				N,J,M	4.0e1
Tetradecane			444 444	N,J,M	1.3e1
Hexadecane				N,J,M	2.7e0
Pentane, 3-ethyl-					
Cyclopentane, 1,2-dimethyl-					
- j - p - marroy rje amioarji					

Table B-9. 0031-END-1.	gr	ASS FLOW RATE ams per second	!		
Project:	01-1062-01-0866	Lab Report Date:	08/20/01		
Run Date:	6/20/2001	Lab Report Status:	Final		
Run Identification:	0031-END-1	<del></del>			. T. (-1
Analyte				Rur   Flag	n Total g/sec
Acetone				B	3.0e-5
				<	1.2e-5
Acrylonitrile					1.0e-6
Benzene Bromobenzene				<	5.0e-7
Bromochloromethane			A DESCRIPTION OF THE PROPERTY		6.3e-7
					5.0e-7
Bromodichloromethane				<u>`</u>	7.6e-7
Bromoform				·	9.7e-7
Bromomethane				<,J	3.7e-6
2-Butanone				<,J	
n-Butylbenzene				< <	6.3e-7 3.6e-7
sec-Butylbenzene				<u>-</u>	
tert-Butylbenzene				l	5.9e-7
Carbon disulfide				<	5.0e-6
Carbon tetrachloride				<u> </u>	6.7e-7
Chlorobenzene				<,J	4.0e-7
Chlorodibromomethane				<	6.3e-7
Chloroethane				<,J	8.4e-7
Chloroform			AAP-W	<b></b> <	1.7e-6
Chloromethane	DAMAGE IN THE THE THE THE			<	1.2e-5
2-Chlorotoluene				<	2.5e-7
4-Chlorotoluene		****		<	2.5e-7
1,2-Dibromo-3-chloropropane				<	1.2e-6
1,2-Dibromoethane				<	8.4e-7
Dibromomethane			- Aller Part Tarre	_ <	7.2e-7
1,2-Dichlorobenzene				<	7.6e-7
1,3-Dichlorobenzene				<	4.2e-7
1,4-Dichlorobenzene	, ,			<	5.9e-7
Dichlorodifluoromethane	A189			<,J	6.7e-7
1,1-Dichloroethane				<	6.3e-7
1,2-Dichloroethane		MARATE TO THE PARTY OF THE PART		<,J	6.7e-7
1,1-Dichloroethene			40.00	<,J	7.6e-7
cis-1,2-Dichloroethene			and the state of t	<	6.3e-7
trans-1,2-Dichloroethene				<	6.7e-7
1,2-Dichloropropane				<,J	5.5e-7
1,3-Dichloropropane			A SANSKI - V	<	7.2e-7
2,2-Dichloropropane				<	6.7e-7
1,1-Dichloropropene				<	7.6e-7
cis-1,3-Dichloropropene		AND THE PROPERTY OF THE PROPER		<	5.5e-7
trans-1,3-Dichloropropene				<	6.3e-7
Ethylbenzene				<	3.9e-7
Hexachlorobutadiene				<	9.3e-7
2-Hexanone				<	2.4e-6
Isopropylbenzene				<	2.9e-7
p-Isopropyltoluene				<	4.6e-7
Methylene chloride				В	7.2e-6
4-Methyl-2-pentanone	12002374			<	2.5e-6
Naphthalene		- ALLONDO TOTAL PORTO		<	8.8e-7

Table B-9. 0031-END-1.		
MASS FLOW RATE		
Project: 01-1062-01-086\( Lab \) Report Date: 0	8/20/01	
Run Date: 6/20/2001 Lab Report Status:	Final	
Run Identification: 0031-END-1		
Analyte	Rur	n Total
	Flag	g/sec
n-Propylbenzene	<	2.8e-7
Styrene	<	3.3e-7
1,1,1,2-Tetrachloroethane	<	4.2e-7
1,1,2,2-Tetrachloroethane	<	9.3e-7
Tetrachloroethene	<	6.3e-7
Toluene	<,J	1.2e-6
1,2,3-Trichlorobenzene	<	8.8e-7
1,2,4-Trichlorobenzene	<	9.3e-7
1,1,1-Trichloroethane	<	8.0e-7
1,1,2-Trichloroethane	<	7.2e-7
Trichloroethene	<	6.7e-7
Trichlorofluoromethane	<,J	6.7e-7
1,2,3-Trichloropropane	<	1.0e-6
1,2,4-Trimethylbenzene	<	4.2e-7
1,3,5-Trimethylbenzene	<	2.5e-7
Vinyl chloride		7.6e-7
m-Xylene & p-Xylene	<	1.9e-6
o-Xylene	<,J	3.3e-7
TICs		
Pentane, 3,3-dimethyl-	N,J,M	2.5e-7
Hexane, 2-methyl-	N,J,M	1.3e-6
Pentane, 2,3-dimethyl-	N,J,M	1.3e-7
Hexane, 3-methyl-	N,J,M	3.2e-6
Cyclohexene	N,J,M	1.5e-7
Cyclobutane, ethenyl-	N,J,M	5.0e-7
Cyclopentane, 1,2-dimethyl-, t	N,J,M	3.3e-7
Cyclohexane, methyl-	N,J,M	8.4e-7
Hexane, 2,4-dimethyl-	N,J,M	7.6e-7
Cyclopentane, ethyl-	N,J,M	1.7e-7
Methane, trichloronitro-	N,J,M	1.5e-6
Benzonitrile	N,J,M	8.0e-7
Undecane	N,J,M	2.1e-6
Undecane, 5-methyl-	N,J,M	7.6e-7
Dodecane	N,J,M	1.1e-4
Tridecane	N,J,M	1.3e-5
Tetradecane	N,J,M	4.2e-6
Hexadecane	N,J,M	8.8e-7
Pentane, 3-ethyl-		
Cyclopentane, 1,2-dimethyl-		· · · · · · · · · · · · · · · · · · ·

Size	Table B-7.	0031-STK	1-2.	VO	ST SAM	PLING DAT	TA SHEF	T	1		
Project   01-1062-01-0866	Site	HLLWE Offe	nas Tie-in		OT OAIII				No.:		2
Date   Gol2   Vol										1	
VOST Tube   Leak Check   Sampling   Sampling   Sampling   Sampling   Sampling   Sampling   Temp.   T											
Sample   Note   Sample   Not				,	<u> </u>			4.7.4			
Numbers   Curimin   (n. Hg)   Curimin   (24 hr)   (min.)   (**C)   (	VOST Tube	Leak C	Check	Sampling	Sa	mpling	Probe	Condenser	Meter	Meter	Pump
Set 1	Sample	Rate (	@ vacuum	Rate						Volume	Vacuum
A-3379	Numbers			(L/min)	(24 hr)	(min.)	(°C)		(°C)	. ,	(in. Hg)
A-3380	Set 1		est	Target:							6.0
A-3381		0.008	6	1							6.0
Set 2											5.0
	A-3381										5.0
Residence	ľ										5.0
Residence   Res											5.0
Residence											5.0 5.0
Post-test   Pretest   Pr						4					5.0
Post-test   9:05   50   130   9:0   26   21:274   10:007   6   9:12   57   130   9:0   26   23:388   10:007   6   9:12   57   130   9:0   26   24:311   10:008   10:00   22   3:46   10:008   10:00   22   3:46   10:008   10:00   22   3:46   10:008   10:008   10:00   22   3:46   10:008   10:008   10:00   22   3:46   10:008   1		·									5.0
Post-test   Q-007   6					·						5.0
Note		Post-	tact								5.0
Total											5.0
Set 2	İ	1-		Total							Max:
Set 2							130	8.1	23	l i	6.0
A-3382 A-3383 A-3384 A-3385 A-3387 A-3484 A-3486 A-3484 A	Set 2	Pret	est		9:27	0	1			0.000	6.0
A-3384				1							6.0
A-3384  A-3384  A-3384  A-3384  A-3384  A-3384  A-3384  A-3384  A-3386  A-3387  A-3387  A-3387  A-3388  A-3387  A-3388  A-3389  A-3380  A-3389  A-388  A-388  B-0003  B-0008									22		6.0
Post-test	A-3384	•			9:42	15	130	11.0	24	9.148	4.0
Post-test						20	130				4.5
Post-test											4.5
Post-test	ļ									4	4.5
Note											4.5
Total										1	4.5
New Pretest   Target:   10:19   0   130   11.2   25		0.000	15				130	11.0	27	<del></del>	4.5
Set 3					0:41	41				24.060	Max:
A-3385 A-3386 A-3387 A-3387 A-3387 A-3387 A-3387 A-3387 A-3388 A-3387 A-3387 A-3387 A-3387 A-3388 A-3387 A-3387 A-3387 A-3388 A-3387 A-3387 A-3388 A-3387 A-3387 A-3387 A-3388 A-3387 A-3388 A-3388 A-3388 A-34 A-348 A-348 A-3488											6.0
A-3386 A-3387 A-3387 A-3387 A-3388 A-3387 A-3388 A-3389 A-340 A-34				Target:							5.0
A-3387  A-3387    10:34		0.006	15								5.0
10:39   20   130   10.0   31   10.751     10:44   25   130   10.0   31   12.879     10:49   30   130   10.0   31   19.033     Post-test   10:59   40   130   11.0   32   21.853     0.006   15   11:04   45   130   11.0   32   24.106     Total   0:45   45   35   30   10.0   31   20.04     Average   130   10.2   31     Set 4   Pretest   Target:   11:12   0   130   10.0   31   2.904     A-3388   0.003   15   11:17   5   130   10.0   31   5.259     A-3389   11:27   15   130   10.0   31   5.259     A-3390   11:37   25   130   10.0   31   10.780     11:32   20   130   10.0   31   10.780     11:32   20   130   10.0   31   10.780     11:37   25   130   10.0   31   10.780     11:47   35   130   10.0   31   10.780     11:47   35   130   10.0   31   10.786     Post-test   11:52   40   130   10.0   31   20.786     Post-test   11:52   40   130   10.1   31     Post-test   10.003   15   Total   0:40   40     Average   130   10.1   31     Post-test   10.003   15   Total   0:40   40     Average   130   10.1   31     Post-test   10.003   15   Total   0:40   40     Average   130   10.1   31     Post-test   10.003   15   Total   0:40   40     Average   130   10.1   31     Post-test   10.003   10.0   30   0.003     Average   10.003   15   Total   0:40   40     Average   10.003   15   Total   0:40   40     Average   10.003   15   Total   0:40   40     Average   10.003   15   Total   0:40   40     Average   10.003   15   Total   0:40   40     Average   10.003   15   10.003   10.003   10.003     A mL		ļ									5.0 5.0
10:44   25   130   10.0   31   12.879	A-3387										5.0
10:49   30   130   10.0   31   16.204     10:54   35   130   10.0   31   19.033     Post-test   10:59   40   130   11.0   32   21.853     0.006   15   11:04   45   130   11.0   32   24.106     Total   0:45   45   130   10.2   31     Average   130   10.2   31     A-3388   0.003   15   11:17   5   130   10.0   31   2.904     A-3389   15   11:22   10   130   10.0   31   2.904     A-3389   11:22   10   130   10.0   31   2.904     A-3389   11:27   15   130   10.0   31   7.951     A-3390   11:27   15   130   10.0   31   10.780     A-3390   11:37   25   130   10.0   31   13.986     A-3391   11:42   30   130   10.0   31   16.832     A-3391   Average   11:52   40   130   11.0   32   24.240     Average   130   10.1   31   20.786     Average   130   10.1   31     Average   130   10.1		-		l							5.0
No.006				<u> </u>							5.0
Post-test											5.0
Na		Post-	test								5.0
Total										I I	5.0
Set 4				Total							Max:
Set 4							130	10.2	31		5.0
A-3388	Set 4	Prete	est		11:12	0			-	0.000	5.0
A-3390							130	10.0	31	2.904	5.0
A-3390							130				5.0
11:37   25   130   10.0   31   13.986					11:27	15		10.0		7.951	5.0
11:42   30   130   10.0   31   16.832     11:47   35   130   10.0   31   20.786     Post-test   11:52   40   130   11.0   32   24.240     0.003   15   Total   0:40   40     24.240   N     Average   130   10.1   31     Condensate   Tenax Rinse: Anasorb Rinse:     A-3391   NA   NA   NA     Final Condensate Volume: Anasorb Rinse Vol.:   NA   mL     40 mL   NA   mL   NA   mL     ADDITIONAL INPUTS     Symbol   Units   Set 1   Set 2   Set 3   Set 4   Average Dry Oxygen Concentration   Co2   %   20.5   20.5   20.5     Process Gas Flow (dry, STP) @ 68°F   Qsd   dscm/sec   0.330   0.330   0.330     CALCULATED SAMPLING PARAMETERS     Symbol   Units   Set 1   Set 2   Set 3   Set 4   Is     Sample Volume @ Standard Conditions   VmStd   dsL   20.383   20.059   19.662   19.777     VmStd=17.647 * Y * Pbar * Vm/(Fm + 460)											5.0
NA											5.0
Post-test   0.003   15   Total   0:40   40   130   11.0   32   24.240   Na   130   10.1   31   24.240   Na   130   10.1   31   24.240   Na   130   10.1   31   24.240   Na   130   10.1   31   24.240   Na   130   10.1   31   24.240   Na   130   10.1   31   24.240   Na   130   10.1   31   24.240   Na   24.240											5.0
O.003   15   Total   O:40   40   130   10.1   31   O:40   Mayerage   O:40   Average   O:40   Average   O:40   O:											5.0
Average							130	11.0	32		5.0
Condensate		0.003	15		0:40	40	400			24.240	Max:
A-3391		· · · · · · · · · · · · · · · · · · ·		Average					31		5.0
Symbol   Units   Set 1   Set 2   Set 3   Set 4   Av	A-3391 Final Conde	nsate Volum	ie:		NA Tenax R	inse Vol.:	NA Anasorb	Rinse Vol.:			j.
Average Dry Oxygen Concentration   Co2					ADDIT	IONAL INP	UTS				
Process Gas Flow (dry, STP) @ 68°F   Qsd   dscm/sec   0.330											Average
CALCULATED SAMPLING PARAMETERS           Symbol         Units         Set 1         Set 2         Set 3         Set 4         I           Sample Volume @ Standard Conditions         VmStd         dsL         20.383         20.059         19.662         19.777           VmStd=17.647 * Y * Pbar * Vm/(Fm + 460)         VmStd         VmS				on							20.5
Symbol Units   Set 1   Set 2   Set 3   Set 4   I	Process Gas F	low (dry, STP	) @ 68°F						0.330	0.330	0.330
Sample Volume @ Standard Conditions   VmStd   dsL   20.383   20.059   19.662   19.777   VmStd=17.647 * Y * Pbar * Vm/(Fm + 460)				CALCUL							
VmStd=17.647 * Y * Pbar * Vm/(Fm + 460)											Net
					VmStd	dsL	20.383	20.059	19.662	19.777	79.881
A A. t I O I De te O A factoria						1 1	C 10-	0.50-	0.500	0.000	0.500
Avg. Actual Sampling Rate, Qm=Vm/min Qm L/min 0.427 0.587 0.536 0.606 Avg. Sampling Rate, QmStd=VmStd/min QmStd dsL/min 0.358 0.489 0.437 0.494											0.539 0.445

CC	NCENTRATIONS			
micrograms p	er dry standard c			
	Lab Report Status	: Final		
0031-STRT-2			Pu	n Total
			E .	μg/dscm
				6.3e1
			< .	3.5e1
				2.8e0
				1.5e0
				1.9e0
				1.5e0
-				2.3e0
				1.9e0
				1.1e1
				1.9e0
				1.1e0
		All a V II W W		1.8e0
				1.6e1
				2.0e0
		3000 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1		1.2e0
				1.9e0
				2.1e0
				4.1e0
				1.5e1
				7.4e-1
		4 - 100000 577		7.4e-1
				3.5e0
				2.5e0
				2.1e0
		A - AN I MINE		2.3e0
				1.3e0
				1.8e0
				2.5e0
				1.9e0 2.0e0
				2.0e0 2.0e0
				1.9e0
				2.0e0
				1.6e0
				2.1e0
				2.1e0 2.0e0
				2.3e0
				1.6e0
				1.9e0
				1.2e0
				2.8e0
				7.0e0
				8.8e-1
			<	1.4e0
			<,B	1.4e1 7.4e0
	micrograms p	01-1062-01- Lab Report Date: 6/21/2001 Lab Report Status	micrograms per dry standard cubic meter 01-1062-01- Lab Report Date: 08/20/01 6/21/2001 Lab Report Status: Final	Micrograms   Per dry standard cubic meter   101-1062-01-   Lab Report Date: 08/20/01   1031-STRT-2

Table B-7. 0031-STRT-2.					
	-	ONCENTRATIONS	• •		
	micrograms	per dry standard cubi	ic meter		
Project:	01-1062-01-	Lab Report Date:	08/20/01		
Run Date:	6/21/2001 0031-STRT-2	Lab Report Status:	Final		
Run Identification: Analyte	0031-31K1-2	1		Run	Total
Analyte					μg/dscm
n-Propylbenzene		_		<	8.3e-1
Styrene				<	9.8e-1
1,1,1,2-Tetrachloroethane					1.3e0
1,1,2,2-Tetrachloroethane				<	2.8e0
Tetrachloroethene				<	1.9e0
Toluene				<,J	3.8e0
1,2,3-Trichlorobenzene				<<	2.6e0
1,2,4-Trichlorobenzene			M W T	<	2.8e0
1,1,1-Trichloroethane				<	2.4e0
1,1,2-Trichloroethane				<	2.1e0
Trichloroethene				<	2.0e0
Trichlorofluoromethane			- 444	<b>&lt;</b> ,J	2.0e0
1,2,3-Trichloropropane				<u> </u>	3.0e0
1,2,4-Trimethylbenzene				_<	1.3e0
1,3,5-Trimethylbenzene			***	<	7.4e-1
Vinyl chloride				<,J	1.6e0
m-Xylene & p-Xylene				_ <	5.5e0
o-Xylene				<	9.8e-1
TICs				<u> </u>	
Hexane, 2-methyl-				N,J,M	3.8e0
Pentane, 2,3-dimethyl-				N,J,M	1.4e0
Hexane, 3-methyl-			AND NO.	N,J,M	3.4e0
Pentane, 3-ethyl-			1.00	N,J,M	4.3e-1
Cyclohexene				N,J,M	2.3e0
Cyclopentane, 1,2-dimethyl-				N,J,M	6.6e-1
Cyclohexane, methyl-				N,J,M	2.5e0
Hexane, 2,4-dimethyl-				N,J,M	2.5e0
Cyclopentane, ethyl-				N,J,M	4.5e-1
Benzonitrile				N,J,M	9.0e-1
Tridecane				N,J,M	1.0e0
Undecane				N,J,M	3.0e0
Decane, 2,2,5-trimethyl-			CONTRACTOR OF THE PARTY OF THE	N,J,M	7.5e-1
Undecane, 5-methyl-				N,J,M	2.6e0
Dodecane				N,J,M	2.8e2
Dodecane, 6-methyl-				N,J,M	6.5e-1
Undecane, 2,6-dimethyl-				N,J,M	5.8e-1
Tridecane				N,J,M	3.0e1
Tetradecane				N,J,M	1.2e1
Pentane, 3,3-dimethyl-	*				

Table B-7. 0031-STRT-2.					
		ASS FLOW RATE	; I		
		rams per second	08/20/01		
Project: Run Date:	6/21/2001	6Lab Report Date: Lab Report Status:	Final		
Run Identification:	0031-STRT-2	Lab Report Status.	Tilla		
Analyte				Rur	Total
•				Flag	g/sec
Acetone	- downers			J,B	2.1e-5
Acrylonitrile				<	1.2e-5
Benzene				_ <,J	9.1e-7
Bromobenzene				<	5.0e-7
Bromochloromethane	MARKET TO			<	6.2e-7
Bromodichloromethane				<	5.0e-7
Bromoform				<	7.4e-7
Bromomethane				<,J	6.2e-7
2-Butanone				<,J	3.7e-6
n-Butylbenzene				<	6.2e-7
sec-Butylbenzene				<	3.5e-7
tert-Butylbenzene				<	5.8e-7
Carbon disulfide				<<	5.4e-6
Carbon tetrachloride				<	6.6e-7
Chlorobenzene				<	3.9e-7
Chlorodibromomethane				<	6.2e-7
Chloroethane				<,J	7.0e-7
Chloroform				<	1.4e-6
Chloromethane				<,J	5.0e-6
2-Chlorotoluene				<	2.4e-7
4-Chlorotoluene				<	2.4e-7
1,2-Dibromo-3-chloropropane				<	1.2e-6
1,2-Dibromoethane				<	8.3e-7
Dibromomethane				<	7.0e-7
1,2-Dichlorobenzene				<	7.4e-7
1,3-Dichlorobenzene				<	4.1e-7
1,4-Dichlorobenzene				<	5.8e-7
Dichlorodifluoromethane				<	8.3e-7
1,1-Dichloroethane				<	6.2e-7
1,2-Dichloroethane			- 1 Nov	<	6.6e-7
1,1-Dichloroethene				<,J	6.6 <b>e-</b> 7
cis-1,2-Dichloroethene				<	6.2e-7
trans-1,2-Dichloroethene				<	6.6e-7
1,2-Dichloropropane				<	5.4e-7
1,3-Dichloropropane				<	7.0e-7
2,2-Dichloropropane				<	6.6e-7
1,1-Dichloropropene				<	7.4e-7
cis-1,3-Dichloropropene				<	5.4e-7
trans-1,3-Dichloropropene				<	6.2e-7
Ethylbenzene				<	3.8e-7
Hexachlorobutadiene				<	9.1e-7
2-Hexanone				<	2.3e-6
Isopropylbenzene				<	2.9e-7
p-Isopropyltoluene		1000		<	4.5 <b>e-</b> 7
Methylene chloride				<,B	4.5e-6
4-Methyl-2-pentanone				<	2.4 <b>e-</b> 6
Naphthalene				<	8.7e-7

Table B-7. 0031-STRT-2.			
	MASS FLOW RATE grams per second		
Project:	01-1062-01-086 Lab Report Date: 08/20/01		
Run Date:	6/21/2001 Lab Report Status: Final		
Run Identification:	0031-STRT-2		
Analyte		Run Flag	Total g/sec
n-Propylbenzene		- r lag	2.7e-7
Styrene		<	3.2e-7
1,1,1,2-Tetrachloroethane		<	4.1e-7
1,1,2,2-Tetrachloroethane		<	9.1e-7
Tetrachloroethene		<	6.2e-7
Toluene		<,J	1.2e-6
1,2,3-Trichlorobenzene		<	8.7e-7
1,2,4-Trichlorobenzene		<	9.1e-7
1,1,1-Trichloroethane		<	7.8e-7
1,1,2-Trichloroethane		<	7.0e-7
Trichloroethene		<	6.6e-7
Trichlorofluoromethane		<,J	6.6e-7
1,2,3-Trichloropropane		<	9.9e-7
1,2,4-Trimethylbenzene		<	4.1e-7
1,3,5-Trimethylbenzene		<	2.4e-7
Vinyl chloride		<,J	5.4e-7
m-Xylene & p-Xylene		<	1.8e-6
o-Xylene		<	3.2e-7
TICs			
Hexane, 2-methyl-		N,J,M	1.2e-6
Pentane, 2,3-dimethyl-		N,J,M	4.5e-7
Hexane, 3-methyl-		N,J,M	1.1e-6
Pentane, 3-ethyl-		N,J,M	1.4e-7
Cyclohexene		N,J,M	7.4e-7
Cyclopentane, 1,2-dimethyl-		N,J,M	2.2e-7
Cyclohexane, methyl-		N,J,M	8.3e-7
Hexane, 2,4-dimethyl-		N,J,M	8.3e-7
Cyclopentane, ethyl-		N,J,M_	1.5e-7
Benzonitrile		N,J,M	3.0e-7
Tridecane		N,J,M	3.4e-7
Undecane		N,J,M	9.9e-7
Decane, 2,2,5-trimethyl-	44.40	N,J,M	2.5e-7
Undecane, 5-methyl-		N,J,M	8.7e-7
Dodecane		N,J,M	9.1e-5
Dodecane, 6-methyl-		N,J,M	2.1e-7
Undecane, 2,6-dimethyl-		N,J,M	1.9e-7
Tridecane		N,J,M	9.9e-6
Tetradecane		N,J,M	4.0e-6
Pentane, 3,3-dimethyl-			

Table B-8.	0031-EN	ND-2.								
			VO	ST SAME	LING DAT	A SHEE				
Site:		Offgas Tie-in	Run No.:		0031-E		Meter Box N	lo.:		2
Project:	01-1062-	1000	Run Type:		Te		Y-factor: 1.005			
Date:	21-	Jun-01	Pbar., in. H	g:	25.2	10	Operator: fe/rw			
VOCETI		l. Ob set	I Oii	Car	lina	Drobo	Candanaar	Motor	Meter	Pump
VOST Tube		k Check	Sampling Rate		npling ime	Probe	Condenser Temp.	Meter Temp.	Volume	Vacuum
Sample Numbers	Rate (L/min)	@ vacuum (in. Hg)	(L/min)	(24 hr)	(min.)	Temp.	(°C)	(°C)	(L)	(in. Hg)
Set 1	,	retest	Target:	13:50		130	14.0	36	0.000	5.0
A-3428	0.003	15	1	13:55	5	130	15.0	36	2.913	5.0
A-3429	0.000			14:00	10	130	15.0	36	5.730	4.5
A-3430		1		14:05	15	130	16.0	36	8.442	4.5
				14:10	20	130	17.0	36	11.143	4.5
				14:15	25	130	14.0	36	13.890	4.5
				14:20	30	130	13.0	37	16.475	4.5
				14:25	35	130	13.0	37	19.112	4.5
ł		·		14:30	40	130	12.0	37	21.676 24.121	4.5 4.5
		st-test	Total	14:35 0:45	45 45	130	12.0	37	24.121	Max:
	0.000	15	Average	0.43	40	130	14.1	36	24.12.1	5.0
Set 2	P	retest	Target:	14:45	0	130	14.0	37	0	4.7
A-3431	0.003	15	1 1		5	130	13.0	37	2.884	4.8
A-3432		t	<u> </u>	14:55	10	130	12.0	37	5.986	4.8
A-3433				15:00	15	130	13.0	37	8.994	4.8
				15:05	20	130	14.0	37	12.108	4.8
				15:10	25	130	13.0	38	16.982	4.8
				15:15	30	130	14.0	37	18.303	4.8
		L		15:20	35	130	13.0	37	21.394	4.8
,		st-test	T-4-1	15:25	40 40	130	13.0	37	25.561 25.561	4.8 Max:
	0,000	15	Total Average	0:40	40	130	13.2	37	20.001	4.8
Set 3		retest	Target:	15:34	0	130	15.0	37	0.000	4.8
A-3434	0.000	15	Target.	15:39	5	130	14.0	37	3.251	4.8
A-3435	0.000	15		15:44	10	130	13.0	38	6.003	4.8
A-3436		·		15:49	15	130	13.0	38	9.136	4.8
				15:54	20	130	14.0	38	11.487	4.8
				15:59	25	130	14.0	38	13.953	4.8
				16:04		130	14.0	38	16.616	4.8
		į		16:09	35	130	14.0	38	19.331	4.8
		l		16:14	40	130	14.0	38	22.039	4.8
		st-test	T-4-1	16:18	44	130	14.0	38	24.214 24.214	4.8 Max:
	0.006	15	Total Average	0:44	44	130	13.9	38	24.214	4.8
Set 4	D	retest	Target:	16:26	0	130	15.0	38	0.000	4.0
A-3437	0.006	15	raiget.	16:31	5	130	15.0	38	2.624	4.0
A-3438		10		16:36	10	130	15.0	38	5.801	4.0
A-3439				16:41		130	15.0	38	8.223	4.0
		T		16:46	20	130	15.0	39	10.998	4.0
]				16:51	25	130	15.0	39	13.737	4.0
		!		16:56	30	130	15.0	39	16.501	4.0
1				17:01	35	130	15.0	39	19.272	4.0
		at toot		17:06	40	130	15.0	40 40	22.038 24.227	4.0
	0.006	st-test 15	Total	17:10 0:44	44	130	15.0	40	24.227	Max:
	0.000	10	Average	U.44		130	15.0	39	-7.22	4.0
Condensate			Avelage	Tenax R	inse:		Rinse:		[	
A-3440	•			NA		NA				
Final Conder	nsate Volu	ıme:			inse Vol.:		Rinse Vol.:			
	mL			NA	mL	NA	mL			
					ONAL INP					
				Symbol	Units	Set 1	Set 2	Set 3	Set 4	Average
Average Dry			<u> </u>	Co2	<u>%</u>	20.5		20.5		20.5
Process Gas F	low (dry, S	TP) @ 68°F		Qsd	dscm/sec	0.330		0.330	0.330	0.330
	ŗ		CALCUL		AMPLING F			Ca4 2	Cat 4	Not
Cample Valu	ma @ St-	andard Candi	tions	Symbol VmStd	Units dsL	<b>Set 1</b> 19.344	Set 2 20.452	<b>Set 3</b> 19.331	Set 4 19.280	<b>Net</b> 78.406
Sample Volu VmStd=17.647		unuaru CONGI	110113	viiiolu	usL	13.344	20.402	15.331	19.200	7 0.400
Avg. Actual S			n/min	Qm	L/min	0.536	0.639	0.550	0.551	0.569
Avg. Samplir	na Rate. C	mStd=VmSt	d/min	QmStd	dsL/min	0.430		0.439		0.455
								· · · · · ·		

Table B-8. 0031-END-2.			
CONCENTRATIONS	· ' '		
micrograms per dry standard cubi Project: 01-1062-01- Lab Report Date:	os/20/01		
Run Date: 6/21/2001 Lab Report Status:	Final		
Run Identification: 0031-END-2			
Analyte			Total
		Flag	μ <b>g/dscn</b>
Acetone		<,J,B	4.8e1
Acrylonitrile		<	3.6e1
Benzene		<	2.6e0
Bromobenzene		<	1.5e0
Bromochloromethane		<	1.9e0
Bromodichloromethane			1.5e0
Bromoform		<	2.3e0
Bromomethane		J	2.6e0
2-Butanone		<	1.1e1
n-Butylbenzene		<	1.9e0
sec-Butylbenzene	A STATE OF THE STA	<	1.1e0
tert-Butylbenzene		<	1.8e0
Carbon disulfide		<	8.2e0
Carbon tetrachloride	A A A A A A A A A A A A A A A A A A A	<,J	2.0e0
Chlorobenzene		<,J	1.2e0
Chlorodibromomethane		<	1.9e0
Chloroethane		<,J	2.0e0
Chloroform		<	5.0e0
Chloromethane		<,J	3.2e1
2-Chlorotoluene		<	7.5e-1
4-Chlorotoluene		<	7.5e-1
1,2-Dibromo-3-chloropropane	. , , ,	<	3.4e0
1,2-Dibromoethane		<	2.6e0
Dibromomethane		<	2.2e0
1,2-Dichlorobenzene		<	2.3e0
1,3-Dichlorobenzene		<u> </u>	1.3e0
1,4-Dichlorobenzene		<	1.8e0
Dichlorodifluoromethane		<,J	2.0e0
1,1-Dichloroethane		<	1.9e0
1,2-Dichloroethane		<	2.0e0
1,1-Dichloroethene		<,J	2.0e0
cis-1,2-Dichloroethene		<	1.9e0
trans-1,2-Dichloroethene		<	2.2e0
1,2-Dichloropropane		<	1.7e0
1,3-Dichloropropane		<	2.2e0
2,2-Dichloropropane		<	2.0e0
1,1-Dichloropropene		<	2.3e0
cis-1,3-Dichloropropene		<	1.7e0
trans-1,3-Dichloropropene		<	1.9e0
Ethylbenzene		<	1.2e0
Hexachlorobutadiene		<	2.9e0
2-Hexanone		<	7.1e0
Isopropylbenzene		<	8.9e-1
p-Isopropyltoluene		<	1.4e0
Methylene chloride		<,J,B	5.5e0

Table B-8. 0031-END-2.					
		NCENTRATIONS			
Decision	micrograms p 01-1062-01-	per dry standard cubi Lab Report Date:	ic meter 08/20/01		
Project: Run Date:	6/21/2001	Lab Report Status:	Final		
Run Identification:	0031-END-2	Lab report otatas.	T III GI		
Analyte				Rur	Total
				Flag	μ <b>g/dscm</b>
4-Methyl-2-pentanone				<	7.5e0
Naphthalene				<	2.7e0
n-Propylbenzene				<	8.4e-1
Styrene				<	9.9e-1
1,1,1,2-Tetrachloroethane				<	1.3e0
1,1,2,2-Tetrachloroethane				<	2.9e0
Tetrachloroethene				<	1.9e0
Toluene				<,J	2.3e0
1,2,3-Trichlorobenzene					2.7e0
1,2,4-Trichlorobenzene				<	2.9e0
1,1,1-Trichloroethane				<	2.4e0
1,1,2-Trichloroethane				<	2.2e0
Trichloroethene				<	2.0e0
Trichlorofluoromethane				<,J	2.0e0
1,2,3-Trichloropropane				<	3.1e0
1,2,4-Trimethylbenzene				<	1.3e0
1,3,5-Trimethylbenzene				. <	7.5e-1
Vinyl chloride				<,J	2.2e0
m-Xylene & p-Xylene				<	5.6e0
o-Xylene				<,J	9.8e-1
TICs					
Hexane, 2-methyl-				N,J,M	3.1e0
Pentane, 2,3-dimethyl-				N,J,M	1.8e0
Hexane, 3-methyl-				N,J,M	4.3e0
Pentane, 3-ethyl-				N,J,M	4.1e-1
Cyclohexene				N,J,M	3.3e-1
Cyclopentane, 1,2-dimethyl-, t				N,J,M	3.8e-1
Cyclohexane, methyl-				N,J,M	1.5e0
Hexane, 2,4-dimethyl-				N,J,M	1.2e0
Benzonitrile				N,J,M	6.0e-1
Undecane				N,J,M	1.5e0
Undecane, 5-methyl-	·			N,J,M	2.3e0
Dodecane				N,J,M	2.2e2
Undecane, 2,6-dimethyl-				N,J,M	1.1e0
Tridecane				N,J,M	4.0e1
Tetradecane				N,J,M	1.8e1
Cyclopentane, ethyl-					
Pentane, 3,3-dimethyl-	· · · · · · · · · · · · · · · · · · ·				

Table B-8. 0031-END-2.				
1 aut D-0. 0031-END-2.	MASS FLOW RATE			
	grams per second			
	-0866 Lab Report Date:	08/20/01		
Run Date: 6/21/200		Final		
Run Identification: 0031-END	J-Z		D	Total
Analyte			Flag	g/sec
Acetone			<,J,B	1.6e-5
Acrylonitrile			\ \ \ \ \ \ \	1.2e-5
Benzene		<del></del>	\ \ <	8.4e-7
Bromobenzene			<	5.0e-7
Bromochloromethane			<	6.3e-7
Bromodichloromethane			<	5.0e-7
Bromoform				7.6e-7
Bromomethane			J	8.4e-7
2-Butanone			<	3.7e-6
n-Butylbenzene			<	6.3e-7
sec-Butylbenzene		m raini samai - wa	<	3.6e-7
tert-Butylbenzene			<	5.9e-7
Carbon disulfide			<	2.7e-6
Carbon tetrachloride		AND DESCRIPTION OF THE PARTY OF	<,J	6.7e-7
Chlorobenzene			-, <u>J</u>	4.0e-7
Chlorodibromomethane	1. 1.		<	6.3e-7
Chloroethane		A BULL THE THE TOTAL OF THE TOT	-,J	6.7e-7
Chloroform			<	1.6e-6
Chloromethane			<,J	1.1e-5
2-Chlorotoluene			<,0	2.5e-7
4-Chlorotoluene				2.5e-7
1,2-Dibromo-3-chloropropane			<	1.1e-6
1,2-Dibromoethane				8.4e-7
Dibromomethane			<	7.1e-7
1,2-Dichlorobenzene				7.6e-7
1,3-Dichlorobenzene	The state of the s		<	4.2e-7
1,4-Dichlorobenzene	MANUAL (1977)		-	5.9e-7
Dichlorodifluoromethane			-,J	6.7e-7
1,1-Dichloroethane			<	6.3e-7
1,2-Dichloroethane				6.7e-7
1,1-Dichloroethene	N		-,J	6.7e-7
cis-1,2-Dichloroethene				6.3e-7
trans-1,2-Dichloroethene			\ \ \ <	7.1e-7
1,2-Dichloropropane		The second secon		5.5e-7
1,3-Dichloropropane	MALAN STARTER STARTER			7.1e-7
2,2-Dichloropropane			<	6.7e-7
• • •			<	7.6e-7
1,1-Dichloropropene			<	5.5e-7
cis-1,3-Dichloropropene				6.3e-7
trans-1,3-Dichloropropene				3.9e-7
Ethylbenzene				
Hexachlorobutadiene	, and the second second			9.7e-7
2-Hexanone				2.4e-6
Isopropylbenzene			<	2.9e-7
p-Isopropyltoluene			<	4.6e-7
Methylene chloride			<,J,B	1.8e-6

Table B-8. 0031-END-2.			
	MASS FLOW RATE		
Project:	grams per second 01-1062-01-086\( Lab \) Report Date: 08/20/01		
Run Date:	6/21/2001 Lab Report Status: Final		
Run Identification:	0031-END-2		
Analyte			Total
A Marthall O and a second		Flag	g/sec
4-Methyl-2-pentanone	er e	<	2.5e-6 8.8e-7
Naphthalene			2.8e-7
n-Propylbenzene Styrene			3.3e-7
1,1,1,2-Tetrachloroethane			4.2e-7
1,1,2,2-Tetrachloroethane		_ <	9.7e-7
Tetrachloroethene		<	6.3e-7
Toluene		+ `,J	7.6e-7
1,2,3-Trichlorobenzene		\ \ \ <	8.8e-7
1,2,4-Trichlorobenzene		<del>  `</del>	9.7e-7
1,1,1-Trichloroethane		-	8.0e-7
1,1,2-Trichloroethane		<del>  `</del>	7.1e-7
Trichloroethene	The second secon	-	6.7e-7
Trichlorofluoromethane		-,J	6.7e-7
1,2,3-Trichloropropane		<	1.0e-6
1,2,4-Trimethylbenzene		<	4.2e-7
1,3,5-Trimethylbenzene		<	2.5e-7
Vinyl chloride		<,J	7.1e-7
m-Xylene & p-Xylene		<	1.9e-6
o-Xylene		<,J	3.2e-7
TICs			
Hexane, 2-methyl-		N,J,M	1.0e-6
Pentane, 2,3-dimethyl-		N,J,M	5.9e-7
Hexane, 3-methyl-		N,J,M	1.4e-6
Pentane, 3-ethyl-		N,J,M	1.3e-7
Cyclohexene		N,J,M	1.1e-7
Cyclopentane, 1,2-dimethyl-,	t	N,J,M	1.3e-7
Cyclohexane, methyl-		N,J,M	5.0e-7
Hexane, 2,4-dimethyl-		N,J,M	4.0e-7
Benzonitrile		N,J,M	2.0e-7
Undecane	· · · · · · · · · · · · · · · · · · ·	N,J,M	5.0e-7
Undecane, 5-methyl-		N,J,M	7.6e-7
Dodecane		N,J,M	7.1e-5
Undecane, 2,6-dimethyl-		N,J,M	3.5e-7
Tridecane		N,J,M	1.3e-5
Tetradecane		N,J,M	5.9e-6
Cyclopentane, ethyl-	The state of the s		
Pentane, 3,3-dimethyl-		<u></u>	

Table B-9. 0050-STRT-1.

## 0050 SAMPLING DATA SHEET FOR HLLWE TESTS

Site:	HLLWE	Offgas Tie-in		Sampling	Location:	-	MAN	N-OFG-73	Nozzle No	.:		2-01	Est. ΔP:	0.15	Est. Tstack, °F:	255
Project:	01-	1062-01-0866		Duct ID, i	nches:			12	Nozzle Size, in.: 0.3140			Est. K:	6.36	Est. vs, ft/s:	28.4	
Date:		6/7/2001		Static Pre	essure, in.	WG:	G: -17.5			Pitot No.: JM-2			Est. ∆H:	0.95	Operator(s): FE,	JA,RW
Run No.:	0	050-STRT-1		Est. O2, 9	<u>%:</u>			Pitot Coeff.: 0.84			0.84	Est. DGM Temperature, °F 80				
Run Type:		TEST		Est CO2,				0	Meter Box	No.		2	Meter Box	Leak Ched		
Phar., in. H		25.200		Est. Mois	<del></del>				ΔH@:				Pretest	0.010	cfm @ 15 in.	. Hg
Tambient, °F		70		Impinger				9	Y-factor:		3.	1.0328			Pitot:	
DGM vol. (	30ai (m*):	3.00		DGM Vol.	Goal (ft <sup>3</sup> )	):		127.080	Min. endin	g DGM vol.	. (ft°):	819.905	Post-test	0.010	cfm @ 11 in.	. Hg
Sampling	Clock	Velocity ∆P		Meter	Actual	Meter			TEMPERA	<del>``</del>			Pump	%I,	00111151170	
Time (min.)	Time (24hr)	ΔΡ (in. WG)		ΔH (in. WG)	ΔH (in, WG)	Volume (cubic feet)	Probe (if heated)	Stack	In Me	ter Out	Filter	lmpinger Exit	Vacuum (in. Hg)	701 <sub>i</sub>	COMMENTS	
0	8:00	0.170	0.412	1.27		692.825	252	133	70	61	258	52	9.1	-		-
10	8:10	0.170	0.412	1.28		699.423	251	134	78	64	258	45	9.2	92		
20	8:20	0.170	0.412	1.29		706.640	250	134	80	67	259	44	9.6	100		
30	8:30	0.160	0.400	1.21		713.400	252	134	. 81	68	259	44	10.1	96		
40	8:40	0.160	0.400	1.22		720.600	252	134	82	69	258	45	10.1	102		
50	8:50	0.160	0.400	1.22		727.830	253	134	83	70	258	45	10.1	103		
60	9:00	0.160	0.400	1.22		735.060	253	134	84	71	259	46	10.1	103		
70	9:10	0.160	0.400	1.22		742.270	249	134	84	72	258	47	10.1	102		
80	9:20	0.160	0.400	1.23		749.510	249	133	85	73	259	47	10.1	102		
90	9:30	0.160	0.400	1.23		756.760	249	133	86	74	258	48	10.1	102		
100	9:40	0.160	0.400	1.23		764.010	249	133	86	74	258	49	10.1	102		
110	9:50	0.160	0.400	1.23		771.270	252	133	88	75	258	49	10.1	102		
120	10:00	0.160	0.400	1.23		778.480	250	133	89	75	258	49	10.1	101		
130	10:10	0.160	0.400	1.23		785.690	251	133	89	77	259	51	10.1	101		
140	10:20	0.160	0.400	1.24		793.120	253	133	90	77	259	51	10.1	104		
150	10:30	0.160	0.400	1.24		800.398	255	133	90	78	258	52	10.1	102		
160	10:40	0.160	0.400	1.24		807.695	248	133	91	79	261	52	10.1	102		
170	10:50	0.160	0.400	1.24		814.890	252	133	91	79	258	53	10.1	101		
180	11:00	0.160	0.400	1.24		822.292	251	133	91	79	258	53	10.1	103		
Total	Total	ΔPavg		Average		Total		Ā١	erage Tem	peratures (	°F)		Max.	Ave. %I		
180	3:00	0.162	0.402	1.24		129.467	251	133	85	73	258	49	10.1	101		

Site:	HLLWE Offg	as Tie-in		Impinger Box no.:	2	9			
Date:	6/7/2001								
Run No.:	0050-STRT-1								_
Component:	KO-1	lmp-1	lmp-2	lmp-3	Imp-4		Acid Scrub Section	on	
Туре:	short stem		G-S	mod	dified	short stem	modified	modified	
Reagent: None  Nominal Contents: Empty		0.0	<b>5M</b> H₂SO₄	0.1N	NaOH	None	<b>2N</b> NaOH	Silica Gel	
			100 mL	100	) mL	Empty	<b>200</b> mL	300-400g	
Post-test Wt., g:	559.8	662.0	669.7	719.5	680.7	582.2	694.2	791.5	Train Wt. Gain
Pre-test Wt., g:	559.9	665.3	667.4	720.1	680.7	566.7	709.4	766.3	Train VVI. Gain
Wt. Gain, g:	-0.1	-3.3	2.3	-0.6	0.0	15.5	-15.2	25.2	23.8
Post-test Volume:	0	100	100	100	100	0			Train Vol. Gain
Pre-test Volume:	0	100	100	100	100	0			Train voi. Gain
Volume Gain:	0	0	0	0	0	0			0.0
Post-test pH:				12.0	13.0		13.0		
									•
		Filter Lot #	: T4208E	H <sub>2</sub> SO <sub>4</sub> Lot #:	32060	NaOH Lot #:	QCLAB-381	DI Water* Lot #:	QCLAB-01
								 * used to dilute acid	and caustic

Record impinger change-out and other important information below:

20.5

0

O2%

CO2%

Table B-9. 0050-STRT-1.

Project:	2-01-0866		
Run Date: Run Identification:		/2001 STRT-1	
PARAMETER	SYMBOL	UNITS	
Absolute Pressure in the Duct	Pabs	in. Hg	23.913
Average Duct Gas Temperature	Ts	R	593
Average Meter Temperature	Tm	R	539
Average Gas Oxygen Content	Co2,m	%	20.5
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0
Total Impinger Weight Gain (water)	Ww	grams	23.8
Nozzle Area	An	ft²	0.000538
Duct Area	As	ft²	0.7854
Sample Volume	VmStd	dscf	110.729
Sample Volume (SI)	VmStdm	dscm	3.136
Average Sampling Rate	Qm	dscf/m	0.615
Volume of Water Vapor	VwStd	scf	1.122
Volume of Water Vapor (SI)	VwStdm	scm	0.0318
Moisture Fraction	Bws	• • • • • • • • • • • • • • • • • • •	0.010
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.71
Gas Velocity at Nozzle	vn	ft/s	26.8
Gas Velocity at Nozzle (SI)	vnm	m/s	8.18
Average Gas Velocity	vncor	ft/s	22.64
Dry Offgas Flow Rate	Qsd	dscf/h	45,066
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,276
Actual Offgas Flow Rate	Q	acf/h	64,013
Intermediate Isokinetic Rate	li	%	101.3
Final Isokinetic Rate	- I	%	101.0

Table B-9. 0050-STRT-1.

Project: 01-1062-01-0866

Run Date: 6/7/2001 Run Identification: 0050-STRT-1

Run Type:

TEST 9/11/2001

Lab Report Date: Lab Report Status:

Final

### 0050 RESULTS

without blank corrections

(preliminary or final)		Final										
		CO	NC	ENTRATION				MA	S	FLOW RAT	ES	3
		Actual (mg/acm)		Standard (mg/scm)		Ory Standard (mg/dscm)		mg/min		grams/sec		lb/h
Vapor Phase Species:		mg/acm		mg/scm		mg/dscm						
Chloride (as HCl)	В	6.5e-1	В	9.2e-1	B B	ppmv	В	2.0e1	В	3.3e-4	В	2.6e-3
Chloride (as Cl2)	<	mg/acm 6.5e-2	<	mg/scm 9.2e-2	V V	ppmv		2.0e0	<	3.3e-5	<	2.6e-4
Fluoride	<	mg/acm 9.0e-2	<	mg/scm 1.3e-1	<	mg/dscm 1.3e-1	٧	2.7e0	<	4.5e-5	<	3.6e-4
Nitrate		mg/acm 2.7e0		mg/scm 3.8e0		mg/dscm 3.8e0		8.1e1		1.4e-3		1.1e-2
Nitrite	<	mg/acm 3.6e-1	<	mg/scm 5.1e-1	<	mg/dscm 5.1e-1	<	1.1e1	<	1.8e-4	<	1.4e-3
Total Particulate							В	2.4e1	В	4.0e-4	В	3.1e-3

Table B-10. 0050-END-1.

### 0050 SAMPLING DATA SHEET FOR HLLWE TESTS

	te: HLLWE Offgas Tie-in Sampling Location: MAN-OFG-73 Nozzle No.: 2-01 Est. ΔP: 0.15 Est. Tstack. °F: 255																
Site:		Offgas Tie-in		Sampling	Location:		MAN	N-OFG-73	Nozzle No	:		2-01	Est. ∆P:	0.15	Est. Tstack, °F	:	255
Project:	01-1	062-01-0866		Duct ID, i	nches:			12	Nozzle Siz	e, in.:		0.3140	Est. K:		Est. vs, ft/s:		28.5
Date:		6/7/2001			essure, in.	WG:			Pitot No.:				Est. ∆H:		Operator(s):	- 1	FE/RW
Run No.:	0	050-END-1		Est. O <sub>2</sub> , 9		·			Pitot Coeff					Temperatur	•		80
Run Type:		TEST		Est CO <sub>2</sub> ,				0	Meter Box	No.				Leak Che			pass
Phar., in. Ho		25.200		Est. Mois					ΔH@:				Pretest	0.000	cfm @	15 in.	Hg
Tambient, °F DGM vol. 0		3.00		Impinger	Box No.: Goal (ft <sup>3</sup> )			9	Y-factor: Min. endin	a DCM val	/£13\.	1.0328				Pitot:	
								127.080	<u>'</u>		. (IL ).	953.478	Post-test	0.000	cfm @	7.1 in.	нg
Sampling Time	Clock Time	Velocity AP		Meter ΔΗ	Actual ΔΗ	Meter Volume	Probe			TURE (°F)	T	Impinger	Pump Vacuum	%l <sub>i</sub>	CON	MENTS	
(min.)	(24hr)	(in. WG)		(in. WG)	(in, WG)	(cubic feet)	(if heated)	Stack	In In	eter Out	Filter	Exit	(in. Hg)	7014		AIMENTS	- 1
0	14:05	0.150	0.387	1.30		826.398	252	133	95	82	260	49	7.2	-			
10	14:15	0.150	0.387	1.30		833.490	260	133	97	83	260	52	7.2	101			
20	14:25	0.150	0.387	1.30		840.853	254	133	98	84	263	54	7.2	105			
30	14:35	0.150	0.387	1.30		848.132	253	133	99	85	261	51	7.2	104			
40	14:45	0.150	0.387	1.30		855.329	253	133	99	85	263	53	7.2	103			
50	14:55	0.150	0.387	1.30		862.760	254	133	100	87	260	52	7.2	106			
60	15:05	0.150	0.387	1.30		870.115	255	133	99	87	261	53	7.2	105			
70	15:15	0.150	0.387	1.30		877.700	255	133	99	87	262	53	7.2	108			
80	15:25	0.150	0.387	1.30		884.820	253	133	99	87	261	54	7.2	101			
90	15:35	0.150	0.387	1.30		892.187	251	132	100	87	261	54	7.2	105			
100	15:45	0.150	0.387	1.30		899.530	251	132	100	87	261	55	7.2	104			
110	15:55	0.150	0.387	1.30		906.890	251	132	100	87	261	55	7.2	105			
120	16:05	0.150	0.387	1.30		914.230	252	132	101	88	261	55	7.2	104			
130	16:15	0.150	0.387	1.30		921.580	252	132	101	89	261	55	7.2	104			
140	16:25	0.150	0.387	1.30		928.824	252	133	100	88	260	55	7.2	103			
150	16:35	0.150	0.387	1.30		936.310	253	132	101	89	261	56	7.2	106			
160	16:45	0.150	0.387	1.30		943.673	253	132	101	88	257	52	7.2	104			
170	16:55	0.150	0.387	1.30		951.037	254	133	101	89	259	57	7.2	104			
180	17:05	0.150	0.387	1.30		958.412	254	132	101	89	259	57	7.2	104			
190	17:15	0.150	0.387	1.30		965.768	254	132	100	89	259	57	7.2	104			
195	17:20	0.150	0.387	1.30		969.463	254	132	100	88	260	59	7.1	105	END of TEST		
Total	Total	ΔPavg		Average	<u> </u>	Total	<u> </u>	A	erage Tem	peratures (	(°F)		Max.	Ave. %l <sub>i</sub>			
195	3:15	0.150	0.387	1.30		143.065	253	133	100	87	261	54	7.2	104			

			_	9	mpinger Box no.:	ı	ffgas Tie-in	HLLWE O	Site:
				a				6/7/2001	Date:
								0050-END-1	Run No.:
	າ	Acid Scrub Section		lmp-4	lmp-3	lmp-2	lmp-1	KO-1	Component:
	modified	modified	short stem	lified	mod	-S	G	short stem	Гуре:
	Silica Gel	<b>2N</b> NaOH	None	NaOH	0.1N	H₂SO₄	0.05M	None	Reagent:
	300-400g	200 mL	Empty	mL	100	) mL	al Contents: Empty 10		Nominal Contents:
<b>-</b>	792.3	689.8	568.9	702.8	695.4	670.5	660.2	560.0	Post-test Wt., g:
Train Wt. Gain	762.0	693.7	564.1	718.2	681.0	668.5	666.5	559.8	Pre-test Wt., g:
26	30.3	-3.9	4.8	-15.4	14.4	2.0	-6.3	0.2	Wt. Gain, g:
			0.0	100.0	100.0	100.0	100.0	NA	Post-test Volume:
Train Vol. Gair			0.0	100.0	100.0	100.0	100.0	NA	Pre-test Volume:
20000000000000000000000000000000000000			0.0	0.0	0.0	0.0	0.0	0.0	Volume Gain:
		12.0	<b>-</b>	12.0	13.0				Post-test pH:

	Filter Lot #: T4208E	H <sub>2</sub> SO <sub>4</sub> Lot #:	328060 NaOH Lot#	000381	DI Water* Lot #: QCLAB1
					* used to dilute acid and caustic
02%	20.5				
CO2%	0				

Record impinger change-out and other important information below:

Table B-10. 0050-END-1.

Project: 01-1062-01-0866 Run Date: 6/7/2001											
Run Date: Run Identification:	0050-										
PARAMETER	SYMBOL										
Absolute Pressure in the Duct	Pabs	in. Hg	23.913								
Average Duct Gas Temperature	Ts	R	592								
Average Meter Temperature	Tm	R	553								
Average Gas Oxygen Content	Co2,m	%	20.5								
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0								
Total Impinger Weight Gain (water)	Ww	grams	26.1								
Nozzle Area	An	ft²	0.000538								
Duct Area	As	ft²	0.7854								
Sample Volume	VmStd	dscf	119.218								
Sample Volume (SI)	VmStdm	dscm	3.376								
Average Sampling Rate	Qm	dscf/m	0.611								
Volume of Water Vapor	VwStd	scf	1.231								
Volume of Water Vapor (SI)	VwStdm	scm	0.0348								
Moisture Fraction	Bws	ı	0.010								
Dry Gas Molecular Weight	Md	g/g-mol	28.82								
Wet Gas Molecular Weight	Ms	g/g-mol	28.71								
Gas Velocity at Nozzle	vn	ft/s	25.8								
Gas Velocity at Nozzle (SI)	vnm	m/s	7.87								
Average Gas Velocity	vncor	ft/s	21.80								
Dry Offgas Flow Rate	Qsd	dscf/h	43,448								
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,230								
Actual Offgas Flow Rate	Q	acf/h	61,639								
Intermediate Isokinetic Rate	li	%	104.4								
Final Isokinetic Rate	I	%	104.1								

### Table B-10. 0050-END-1.

Project:

01-1062-01-

0866

6/7/2001 Run Date: Run Identification: 0050-END-1

Run Type:

TEST

Lab Report Date: Lab Report Status: 9/11/2001

(preliminary or final)

Final

### 0050 RESULTS

#### • without blank corrections

(preliminary or linar)	-	CONCENTRATIONS MASS FLOW RATES										
			)N(	CENTRATIO				M.A	155	FLOW RAT	ES	
		Actual		Standard	1	ry Standard	l					
		(mg/acm)		(mg/scm)		(mg/dscm)		mg/min	Ç	grams/sec		lb/h
Vapor Phase		mg/acm		mg/scm		mg/dscm						
Species:		_		-		_	l					
Chloride (as HCl)	В	6.1e-1	В	8.5e-1	В	8.6e-1	В	1.8e1	В	2.9e-4	В	2.3e-3
						ppmv	l					
	_	<del> </del>			В	5.7e-1	上					
Chloride (as Cl2)		mg/acm		mg/scm		mg/dscm	l					
	В	1.2e-1	В	1.7e-1	В	1.7e-1	В	3.5e0	В	5.8e-5	В	4.6e-4
						ppmv	l					
					В	5.7e-2						
Fluoride		mg/acm		mg/scm		mg/dscm						
	<	8.4e-2	<	1.2e-1	<	1.2e-1	<	2.4e0	<	4.0e-5	<	3.2e-4
Nitrate	$\vdash$	mg/acm		mg/scm	-	mg/dscm	╁				-	
		2.2e0		3.0e0		3.1e0		6.3e1		1.1e-3		8.4e-3
Nitrite	<del>                                     </del>	mg/acm		mg/scm	_	mg/dscm	┢		-	***		
	<	7.9e-1	<	1.1e0	<	1.1e0	<	2.3e1	<	3.8e-4	<	3.1e-3
Total Particulate	-					*	-				_	
Total Particulate							В	1.8e1	В	2.9e-4	В	2.3e-3
										,	-	

Table B-11. 0050-STRT-2.

## 0050 SAMPLING DATA SHEET FOR HLLWE TESTS

Site:	HLLWE (	Offgas Tie-in		Sampling	Location	:	IAM	N-OFG-73	Nozzle No	).:		2-01	Est. ∆P:	0.15	Est. Tstack, °F:	133
Project:		062-01-0866		Duct ID,		-		12	Nozzle Si				Est. K:		Est. vs, ft/s:	26.0
Date:		6/11/2001		Static Pre	essure, in	. WG:		-17.5	Pitot No.:	,		<del></del>	Est. ΔH:		Operator(s):	RW, FE
Run No.:	00	50-STRT-2		Est. O <sub>2</sub> ,	%:		•	20.5	Pitot Coef	f.:		0.84	Est. DGM	Temperature, °	<del></del>	80
Run Type:		TEST		Est CO2,	%:		•	0	Meter Box	No.		1	Pitot Leak	Check: pass	X fail	
P <sub>bar.</sub> , in. H		25.009		Est. Mois	st., %:			1.3%	ΔH@:			1.5673	Meter Box	Leak Checks:		
Tambient, °F		65		Impinger				8	Y-factor:				Pretest		16.5 in. Hg time:	7:53
DGM vol.	Goal (m²):	3.00		DGM vol	. Goal (ft)	:		127.080	Min. endir	ng DGM vo	l. (ft):	1096.725	Post-test	0.000 cfm @	7.5 in. Hg time:	1140
Sampling	Clock	Velocity		Meter	Actual	Meter				ATURE (°F)			Pump			
Time (min.)	Time (24hr)	ΔP (in. WG)		ΔH (in. WG)	∆H (in, WG)	Volume (cubic feet)	Probe (if heated)	Stack	ln Me	eter Out	Filter	Impinger Exit	Vacuum (in. Hg)	%I <sub>i</sub>	COMMENTS	
0	7:50	0.150	0.387	1.30		969.645	253	133	63	59	259	51	7.5			
10	8:00	0.150	0.387	1.20		976.751	254	133	64	60	259	58	7.5	107		
20	8:10	0.150	0.387	1.20		983.261	258	133	74	62	259	48	7.5	97		
30	8:20	0.150	0.387	1.20		990.326	259	133	79	63	259	48	7.5	104		
40	8:30	0.150	0.387	1.25		997.109	259	133	78	66	260	50	7.5	100		
52	8:42	0.150	0.387	1.25		1,006.398	259	133	78	68	260	52	7.5	114	Shut down train, operation	ns is having
52	9:20	0.150	0.387	1.25		1,006.398	254	133	76	69	262	51	7.5		instrumentation problems	
62	9:30	0.150	0.387	1.25		1,014.129	257	133	86	71	261	47	7.5	113	Restart train at 0920.	
72	9:40	0.150	0.387	1.25		1,020.596	257	133	87	73	261	51	7.5	94		
82	9:50	0.150	0.387	1.28		1,029.962	257	133	88	74	262	53	7.5	136		
92	10:00	0.150	0.387	1.25		1,035.175	257	133	88	75	261	54	7.5	75		
102	10:10	0.150	0.387	1.25		1,042.451	257	133	89	76	260	54	7.5	105		
112	10:20	0.150	0.387	1.25		1,049.745	258	133	90	77	260	55	7.5	105		
122	10:30	0.150	0.387	1.25		1,056.999	258	133	90	77	262	55	7.5	105		
132	10:40	0.150	0.387	1.25		1,064.621	258	133	90	77	262	55	7.5	110		
142	10:50	0.150	0.387	1.25		1,071.645	254	133	92	79	261	56	7.5	101		
152	11:00	0.150	0.387	1.25		1,078.976	255	133	92	79	<b>2</b> 62	56	7.5	105		
162	11:10	0.150	0.387	1.25		1,086.382	250	133	92	80	260	56	7.5	106		
172	11:20	0.150	0.387	1.25		1,093.731	250	133	92	80	260	57	7.5	106		,
182	11:30	0.150	0.387	1.25		1,100.084	248	133	93	81	261	57	7.5	91		
Total	Total	$\Delta Pavg$		Average		Total		A	erage Ten	peratures	(°F)	1	Max.	Ave. %I <sub>i</sub>		
182	3:02	0.150	0.387	1.25		130.439	256	133	84	72	261	53	7.5	104		

Site:	HLLWE C	ffgas Tie-in	lı	mpinger Box no.:					
Date:	6/11/2001								
Run No.:	0050-STRT-2								
Component:	KO-1	lmp-1	lmp-2	lmp-3	lmp-4		Acid Scrub Section	n	
Туре:	short stem	G	-S	mo	dified	short stem	modified	modified	
Reagent:	None	0.05M	H₂SO₄	0.1N	NaOH	None	2N NaOH	Silica Gel	
ominal Contents: Empty		100	mL	10	0 mL	Empty	<b>200</b> mL	300-400g	
Post-test Wt., g:	571.9	725.0	677.4	698.2	711.1	571.5	706.2	807.5	Train Wt. Gain
Pre-test Wt., g:	571.8	730.6	671.9	699.3	712.9	586.1	706.1	780.0	Train Wt. Gain
Wt. Gain, g:	0.1	-5.6	5.5	-1.1	-1.8	-14.6	0.1	27.5	10.1
Post-test Volume:	0.0	100.0	100.0	100.0	100.0	0.0			Train Vol. Gain
Pre-test Volume:	0.0	100.0	100.0	100.0	100.0	0.0			Train voi. Gain
Volume Gain:	0.0	0.0	0.0	0.0	0.0	0.0			0.0
Post-test pH:				13.0	13.0		14.0		

		Filter Lot #: T4208E	H <sub>2</sub> SO <sub>4</sub> Lot #:	328060	DI Water* Lot #: QCLAB-1
					* used to dilute acid and caustic
O2%	20.5				
CO2%	0				

Record impinger change-out and other important information below:

Table B-11. 0050-STRT-2.

Project:	01-1062	-01-0866	
Run Date:			
Run Identification: PARAMETER	SYMBOL	TRT-2 UNITS	
Absolute Pressure in the Duct	Pabs	in. Hg	23.722
Average Duct Gas Temperature	Ts	R	593
Average Meter Temperature	Tm	R	538
Average Gas Oxygen Content	Co2,m	%	20.5
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0
Total Impinger Weight Gain (water)	Ww	grams	10.1
Nozzie Area	An	ft²	0.000538
Duct Area	As	ft²	0.7854
Sample Volume	VmStd	dscf	110.880
Sample Volume (SI)	VmStdm	dscm	3.140
Average Sampling Rate	Qm	dscf/m	0.609
Volume of Water Vapor	VwStd	scf	0.476
Volume of Water Vapor (SI)	VwStdm	scm	0.0135
Moisture Fraction	Bws	•	0.004
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.77
Gas Velocity at Nozzle	vn	ft/s	25.9
Gas Velocity at Nozzle (SI)	vnm	m/s	7.90
Average Gas Velocity	vncor	ft/s	21.87
Dry Offgas Flow Rate	Qsd	dscf/h	43,468
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,231
Actual Offgas Flow Rate	Q	acf/h	61,842
Intermediate Isokinetic Rate	li	%	104.6
Final Isokinetic Rate	I	%	103.7

## Table B-11. 0050-STRT-2.

01-1062-01-Project:

0866

Run Date: 6/11/2001

Run Identification: 0050-STRT-2 Run Type:

**TEST** 9/11/2001

Lab Report Date: Lab Report Status: (preliminary or final)

Final

### 0050 RESULTS

• without blank corrections

(preliminary or final)	,		WASS FLOW DATES									
	L	cc	)NC	CENTRATIO				MA	SS	FLOW RAT	ES	
		Actual		Standard	D	ry Standard	l					
		(mg/acm)		(mg/scm)		(mg/dscm)		mg/min	Ç	grams/sec		lb/h
Vapor Phase Species:		mg/acm		mg/scm		mg/dscm						
Chloride (as HCl)	В	6.5e-1	В	9.2e-1	В	9.2e-1 ppmv	В	1.9e1	В	3.2e-4	В	2.5e-3
					В	6.1 <u>e-1</u>				·		
Chloride (as Cl2)	٧	mg/acm 6.3e-2	<	mg/scm 8.9e-2		mg/dscm 8.9e-2 ppmv		1.8e0	<	3.0e-5	<	2.4e-4
Clueride	_	malaam	_	mg/scm	<	3.0e-2 mg/dscm	┞		ļ		_	<del></del>
Fluoride	<	mg/acm 8.5e-2	<	1.2e-1	<	1.2e-1	<	2.5e0	<	4.1e-5	<	3.3e-4
Nitrate		mg/acm 1.9e0		mg/scm 2.7e0		mg/dscm 2.7e0		5.6e1		9.4e-4		7.4e-3
Nitrite	<	mg/acm 3.6e-1	<	mg/scm 5.1e-1	<	mg/dscm 5.1e-1	<	1.0e1	<	1.7e-4	<	1.4e-3
Total Particulate								7.2e0		1.2e-4		9.5e-4

Table B-12. 0050-END-2.

## 0050 SAMPLING DATA SHEET FOR HLLWE TESTS

Site:	HLLWE (	Offgas Tie-in		Sampling	Location	:	1AM	N-OFG-73	Nozzle No	D.:		2-01	Est. ∆P:	0.15	Est. Tstack, °F:	133
Project:	01-10	062-01-0866		Duct ID,	inches:		,	12	Nozzle Siz	ze, in.:		0.3140		7.66	Est. vs, ft/s:	26.0
Date:		6/11/2001		Static Pro	essure, in	. WG:		-17.5	Pitot No.:				Est. ∆H:		Operator(s):	FE/RW
Run No.:		050-END-2		Est. O <sub>2</sub> ,				20.5	Pitot Coef					Temperature, °l		80
Run Type:		TEST		Est CO <sub>2</sub> ,				0	Meter Box	No.				Check: pass		PASS
Pbar., in. H		24.975		Est. Mois					ΔH@:					Leak Checks:		
Tambient, °F DGM vol. 0		70 3.00			Box No.: Goal (ft)			9	Y-factor: Min. endir	on DOM voi	(66).		Pretest		15.0 in. Hg time:	13:45
								127.080			. (11).	228.239	Post-test	0 cfm@	8.7 in. Hg time:	17:35
Sampling Time	Clock Time	Velocity ΔP		Meter ΔH	Actual ΔH	Meter Volume	Probe			ATURE (°F) eter		Impinger	Pump Vacuum	%I,	COMMENTS	
(min.)	(24hr)	(in. WG)		(in. WG)	(in, WG)	(cubic feet)	(if heated)	Stack	ln Ivi	Out	Filter	Exit	(in. Hg)	7011	COMMENTS	
0	14:30	0.150	0.387	1.14		101.159	247	133	75	74	260	58	8.2	-		
10	14:40	0.150	0.387	1.15		108.120	248	133	82	75	260	50	8.2	101		
20	14:50	0.150	0.387	1.15		115.120	247	133	86	76	259	52	8.2	101		
30	15:00	0.150	0.387	1.16		122.090	248	132	88	77	259	53	8.2	101		
40	15:10	0.150	0.387	1.16		129.086	248	133	90	78	260	54	8.2	101		
50	15:20	0.150	0.387	1.16		136.050	247	132	91	79	259	54	8.2	100		
60	15:30	0.150	0.387	1.17		143.060	247	132	92	80	260	55	8.2	100		
70	15:40	0.150	0.387	1.17	-	150.050	247	132	92	81	260	55	8.2	100		
80	15:50	0.150	0.387	1.17		157.070	247	132	93	81	259	55	8.2	100		
90	16:00	0.150	0.387	1.17		164.270	247	132	92	81	261	56	8.5	103		
100	16:10	0.150	0.387	1.17		171.589	247	132	93	81	259	57	8.5	105		
110	16:20	0.150	0.387	1.17		178.920	247	132	93	81	260	57	8.7	105		
120	16:30	0.150	0.387	1.17		186.231	247	132	93	82	260	57	8.7	104		
130	16:40	0.150	0.387	1.17		193.623	247	132	93	82	259	58	8.7	106		
140	16:50	0.150	0.387	1.17		201.132	247	132	93	82	259	58	8.7	107		
150	17:00	0.150	0.387	1.17		208.340	247	132	93	82	258	58	8.7	103		
160	17:10	0.150	0.387	1.17		215.732	247	132	93	82	258	58	8.7	106		
170	17:20	0.150	0.387	1.17		222.973	247	132	92	81	260	59	8.7	104		
180	17:30	0.150	0.387	1.17		230.321	247	132	92	81	260	59	8,7	105		
Total	Total	ΔPavg		Average		Total		A\	verage Ten	nperatures	(~F)	T	Max.	Ave. %I <sub>i</sub>		
180	3:00	0.150	0.387	1.16		129.162	247	132	90	80	259	56	8.7	103		

Site:	HLLWE O	ffgas Tie-in		Impinger Box no.:	9	<del></del>			
Date:	6/11/2001								
Run No.:	0050-END-2								
Component:	KO-1	lmp-1	lmp-2	lmp-3	lmp-4		Acid Scrub Section	on -	
Туре:	short stem	G	-S	mod	dified	short stem	modified	modified	
Reagent:	None	0.05M	H₂SO₄	0.1N	NaOH	None	2N NaOH	Silica Gel	_
Nominal Contents:	Empty	100	mL	100	) mL	Empty	200 mL	300-400g	
Post-test Wt., g:	564.0	658.9	668.3	682.5	719.7	592.5	717.3	816.8	T : W
Pre-test Wt., g:	563.9	664.4	666.8	681.7	718.6	589.5	720.0	787.6	Train Wt. Gain
Wt. Gain, g:	0.1	-5.5	1.5	0.8	1.1	3.0	-2.7	29.2	27.5
Post-test Volume:	0.0	100.0	100.0	100.0	100.0	0.0			T : VIO:
Pre-test Volume:	0.0	100.0	100.0	100.0	100.0	0.0			Train Vol. Gain
Volume Gain:	0.0	0.0	0.0	0.0	0.0	0.0			0.0
Post-test pH:				13.0	13.0		14.0		

	Filter Lot #: T408E	H <sub>2</sub> SO <sub>4</sub> Lot #:	328060	NaOH Lot3: 000381	DI Water* Lot #: QCLAB1
	· <del></del>				* used to dilute acid and caustic
02%	20.5				
CO2%	0				

Record impinger change-out and other important information below:

Table B-12. 0050-END-2.

Project:	01-1062	-01-0866	
Run Date: Run Identification:		'2001 END-2	
PARAMETER	SYMBOL		
Absolute Pressure in the Duct	Pabs	in. Hg	23.688
Average Duct Gas Temperature	Ts	R	592
Average Meter Temperature	Tm	R	545
Average Gas Oxygen Content	Co2,m	%	20.5
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0
Total Impinger Weight Gain (water)	Ww	grams	27.5
Nozzle Area	An	ft²	0.000538
Duct Area	As	ft²	0.7854
Sample Volume	VmStd	dscf	108.235
Sample Volume (SI)	VmStdm	dscm	3.065
Average Sampling Rate	Qm	dscf/m	0.601
Volume of Water Vapor	VwStd	scf	1.297
Volume of Water Vapor (SI)	VwStdm	scm	0.0367
Moisture Fraction	Bws		0.012
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.69
Gas Velocity at Nozzle	vn	ft/s	26.0
Gas Velocity at Nozzle (SI)	vnm	m/s	7.91
Average Gas Velocity	vncor	ft/s	21.90
Dry Offgas Flow Rate	Qsd	dscf/h	43,197
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,223
Actual Offgas Flow Rate	Q	acf/h	61,934
Intermediate Isokinetic Rate	li	%	103.1
Final Isokinetic Rate	1	%	103.0

## Table B-12. 0050-END-2.

Project: 01-1062-01-

0866

Run Date: 6/11/2001 Run Identification: 0050-END-2

Run Type:

TEST 9/11/2001

Lab Report Date: Lab Report Status: (preliminary or final)

Final

#### 0050 RESULTS

• without blank corrections

(preliminary or final)													
		CC	N	CENTRATIO				MA	SS	FLOW RAT	ES		
		Actual		Standard	D	ry Standard							
		(mg/acm)		(mg/scm)		(mg/dscm)		mg/min	Ç	grams/sec		lb/h	
Vapor Phase Species:		mg/acm		mg/scm		mg/dscm							
Chloride (as HCl)	В	6.6e-1	В	9.4e-1	ВВ	9.5e-1 ppmv 6.2e-1	В	1.9e1	В	3.2e-4	В	2.6e-3	
Chloride (as Cl2)	<	mg/acm 6.6e-2	<	mg/scm 9.4e-2	<	mg/dscm 9.5e-2 ppmv 3.2e-2		1.9e0	<	3.2e-5	٧	2.6e-4	
Fluoride	<	mg/acm 8.9e-2	٧	mg/scm 1.3e-1	<	mg/dscm 1.3e-1	<	2.6e0	<	4.3e-5	<	3.4e-4	
Nitrate		mg/acm 1.6e0		mg/scm 2.3e0		mg/dscm 2.3e0		4.7e1		7.9e-4		6.2e-3	
Nitrite	<	mg/acm 7.1e-1	<	mg/scm 1.0e0	<	mg/dscm 1.0e0	<	2.1e1	<	3.4e-4	<	2.7e-3	
Total Particulate						anana, .	В	5.3e0	В	8.9e-5	В	7.0e-4	

Table B-13. 0060-STRT-1.

1.20

Average

1.21

217.943

Total

127.667

253

250

133

133

0.150

ΔPavg

0.159

180

Total

180

12:30

Total

3:00

### METHOD 0060 SAMPLING DATA SHEET FOR HLLWE TESTS

Site:	HLLWE O	ffgas Tie-in	Sampling	Location:	MA	N-OFG-73	Nozzie No	).:		2-01	Est. ΔP:	0.17	Est. Tst	ack, °F:		133
Project:		62-01-0866	· · · · · · · · · · · · · · · · · · ·			12	Nozzle Siz				Est. K:		Est. vs,	<del></del>		27.6
Date:		6/5/2001	Static Pre	essure, in. WG:		-17.5	Pitot No.:				Est. ∆H:	1.30	Operate	or(s):	RW	/,FE,JA
Run No		60-STRT-1				20.5	Pitot Coef			0.84	Est. DGM 7	Гетрегаtu	re, °F			80
Run Tyr			Est CO <sub>2</sub> ,		· · · · · · · · · · · · · · · · · · ·	0	Meter Box	No.			Leak Chec					
Pbar., in.			Est. Mois				ΔH@:			1.5673	<del>-</del>	Pre-		Post-		Pass
Tambient,	r: ol. Goal (m³)	3.0		Box No.: . Goal (ft <sup>3</sup> ):		8 127.1	Y-factor:	g DGM vol	/ft <sup>3</sup> \·		DGM Pre:		cfm @			inHg
						127.1			. (IL <i>)</i> .	217.356	DGM Post:	0.000	cfm @		8	inHg
Sampling Time	Clock Time	Velocity ΔP	Meter ∆H	Meter Volume	Heated			TURE (°F)		Impinger	Pump Vacuum	%l <sub>i</sub>		СОММЕ	NTS	
(min.)	(24hr)	(in. WG)	(in. WG)	(cubic feet)	Line	Stack	ln	Out	Filter	Exit	(in. Hg)					
0	9:30	0.160	1.30	90.276	249	133	72	70	260	61	8.0	_	<u></u>			
10	9:40	0.160	1.20	97.240	249	133	80	71	260	53	7.0	99	O2=20.5			
20	9:50	0.160	1.25	104.200	252	134	82	71	263	55	7.5	99				
30	10:00	0.160	1.22	111.220	251	133	84	72	262	57	7.5	99	O2=20.5			
40	10:10	0.160	1.20	118.230	250	133	87	74	260	59	7.5	98	O2=20.6			
50	10:20	0.160	1.20	125.380	249	132	90	77	261	61	7.5	100	O2=20.6			
60	10:30	0.160	1.20	132.315	250	133	89	77	262	58	7.5	97	O2=20.6			
70	10:40	0.160	1.20	139.455	250	133	87	76	261	54	7.5	100	O2=20.6			
80	10:50	0.160	1.20	146.567	250	134	85	74	262	52	7.5	100	O2=20.6			
90	11:00	0.160	1.20	153.653	250	133	81	74	261	52	7.5	100	O2=20.6			
100	11:10	0.160	1.20	160.757	250	133	88	76	262	53	7.5	99	O2=20.6			
110	11:20	0.160	1.20	167.850	251	133	90	77	260	54	7.5	99	O2=20.6			
120	11:30	0.160	1.20	175.250	248	133	91	79	261	54	7.5	103	O2=20.6			
130	11:40	0.160	1.20	182.110	252	133	92	79	260	55	7.5	95	O2=20.6			
140	11:50	0.160	1.20	189.255	249	132	93	81	259	57	7.5	99	O2=20.6			
150	12:00	0.160	1.20	196.425	249	133	94	82	261	57	7.5	99	O2=20.6			
160	12:10	0.160	1.20	203.600	249	133	91	81	260	56	7.5	100	O2=20.6			
170	12:20	0.160	1.20	210.810	250	133	89	79	261	55	7.5	101	O2=20.6			

87

87

Average Temperatures (°F)

77

76

260

261

54

56

7.5

Max.

8.0

O2=20.6

Ave. %I,

Table B-13. 0060-STRT-1.

Site: HLLWE Offgas Tie-in IMPINGER BOX NO. = 6/5/2001 Date: 0060-STRT-1 Run No.: Mercury-Only Section Component: KO-1 Imp-2 Imp-4 Acid Scrub Section Imp-1 Imp-3 Imp-5 Type: short stem modified G-S modified modified short stem modified modified Reagent: None 5% HNO<sub>3</sub> / 10% H<sub>2</sub>O<sub>2</sub> None 4% KMnO<sub>4</sub> / 10% H<sub>2</sub>SO<sub>4</sub> 2N NaOH Silica Gel None Nominal Contents: 100 mL solution in each 200 mL solution in each Empty **Empty Empty** 100 mL 300-400q Post-test Wt., g: 571.9 718.1 704.9 618.6 733.6 731.3 590.0 680.5 900.3 Impinger Pre-test Wt., g: 572.0 720.9 699.5 616.9 731.8 728.8 589.6 686.4 873.2 wt. gain (g) -2.8 Wt. Gain, g: -0.1 5.4 1.7 2.5 -5.9 30.1 1.8 0.4 27.1 Post-test Volume: 0.0 100.0 90.0 0.0 90.0 0.0 100.0 Impinger Pre-test Volume: 0.0 100.0 100.0 0.0 100.0 100.0 0.0 vol. gain (mL) Volume Gain: 0.0 0.0 -10.0 0.0 0.0 -10.0 0.0 -20.0 Post-test pH: 14.0 Filter Lot # H<sub>2</sub>SO<sub>4</sub> Lot # 328060 53322 HNO<sub>3</sub> Lot # 129100 DI water\* Lot # QCLab-1 H<sub>2</sub>O<sub>2</sub> Lot # 992809 KMnO<sub>4</sub> Lot # 006655

* used to dilute the other	er reagents
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 $\frac{O_2\%}{CO_2\%}$  20.6 0.0

Table B-13. 0060-STRT-1.

Project:	01-1062	-01-0866	
Run Date:			
Run Identification:	0060-S SYMBOL		
PARAMETER Absolute Pressure in the Duct	Pabs	in. Hg	23.732
Average Duct Gas Temperature	Ts 	R _	593
Average Meter Temperature	Tm	R	541
Average Gas Oxygen Content	Co2,m	%	20.6
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0
Total Impinger Weight Gain (water)	Ww	grams	30.1
Nozzle Area	An	ft²	0.000538
Duct Area	As	ft²	0.785
Sample Volume	VmStd	dscf	107.88
Sample Volume (SI)	VmStdm	dscm	3.055
Average Sampling Rate	Qm	dscf/m	0.599
Volume of Water Vapor	VwStd	scf	1.419
Volume of Water Vapor (SI)	VwStdm	scm	0.0402
Moisture Fraction	Bws	_	0.013
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.68
Gas Velocity at Nozzle	vn	ft/s	26.8
Gas Velocity at Nozzle (SI)	vnm	m/s	8.16
Average Gas Velocity	vncor	ft/s	22.58
Dry Offgas Flow Rate	Qsd	dscf/h	44,506
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,260
Actual Offgas Flow Rate	Q	acf/h	63,850
Intermediate Isokinetic Rate	li	%	99.6
Final Isokinetic Rate	ı	%	99.6

Table B-13. 0060-STRT-1.

Project: 01-1062-01-0866

6/5/2001 Run Date:

Run Identification: 0060-STRT-1 Run Type:

Test 8/28/2001

Lab Report Date: Lab Report Status: (preliminary or final)

Final

### **RESULTS**

### without blank corrections

(preliminary or linar)												
		CC	DNC	ENTRATIO	NS			M.A	SS F	LOW RAT	ES	
		Actual	S	Standard	Dry	Standard	l					
	(	µg/acm)	(	μg/scm)	(μ	ig/dscm)		μg/min	gr	ams/sec		lb/h
Aluminum (Al)		3.2e1		4.5e1		4.6e1		9.6e2		1.6e-5		1.3e-4
Antimony (Sb)	В	1.0e0	В	1.5e0	В	1.5e0	В	3.2e1	В	5.3e-7	В	4.2e-6
Arsenic (As)	<,B	3.7e-1	<,B	5.2e-1	<,B	5.2e-1	<,B	1.1e1	<,B	1.8e-7	<,B	1.5e-6
Barium (Ba)	В	1.5e0	В	2.2e0	В	2.2e0	В	4.6e1	В	7.7e-7	В	6.1e-6
Beryllium (Be)	<,B	1.5e-1	<,B	2.1e-1	<,B	2.1e-1	<,B	4.5e0	<,B	7.4e-8	<,B	5.9e-7
Cadmium (Cd)	В	1.0e-1	В	1.5e-1	В	1.5e-1	В	3.2e0	В	5.3e-8	В	4.2e-7
Chromium (Cr)		6.8e-1		9.7e-1		9.8e-1		2.1e1		3.4e-7		2.7e-6
Cobalt (Co)	В	6.2e-1	В	8.7e-1	В	8.8e-1	В	1.9e1	В	3.1e-7	В	2.5e-6
Copper (Cu)	В	9.8e-1	В	1.4e0	В	1.4e0	В	3.0e1	В	4.9e-7	В	3.9e-6
Lead (Pb)	<,B	3.0e-1	<,B	4.2e-1	<,B	4.3e-1	<,B	8.9e0	<,B	1.5e-7		1.2e-6
Manganese (Mn)		4.3e0		6.1e0		6.2e0		1.3e2		2.2e-6		1.7e-5
Mercury (Hg)		2.3e1		3.2e1		3.3e1		6.9e2	l .	1.1e-5		9.1e-5
Nickel (Ni)	В	1.1e0	В	1.6e0	В	1.6e0	В	3.4e1	В	5.7e-7	В	4.5e-6
Selenium (Se)	В	7.3e-1	В	1.0e0	В	1.0e0	В	2.2e1	В	3.7e-7	В	2.9e-6
Silver (Ag)	<	5.9e-1	<	8.4e-1	<	8.5e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Thallium (Tl)	<	5.7e-1	<	8.1e-1	<	8.2e-1	<	1.7e1	<	2.9e-7	<	2.3e-6
Vanadium (V)	<	6.2e-1	<	8.7e-1	<	8.8e-1	<	1.9e1	<	3.1e-7	<	2.5e-6
Zinc (Zn)		1.4e1		2.0e1		2.0e1		4.2e2		7.0e-6		5.5e-5

Table B-13. 0060-STRT-1.

Project: 01-1062-01-0866

Run Date:

6/5/2001

Run Type:

Run Identification: 0060-STRT-1

Lab Report Date:

Test 8/28/2001

Lab Report Status:

Final

### **RESULTS**

#### CORRECTED FOR CONTRIBUTION FROM

Corrected for Reagent Blank per EPA SW-846 Test Method 0060

(preliminary or final)												
		CC	NCE	NTRATIO	NS			MA	SS F	LOW RAT	ES	
	7	Actual	St	tandard	Dry	Standard						
	(μ	g/acm)	(h	ıg/scm)	(µ	g/dscm)		µg/min	gr	ams/sec		lb/h
Aluminum (Al)		3.0e1		4.2e1		4.2e1		8.9e2		1.5e-5		1.2e-4
Antimony (Sb)	В	2.4e-1	В	3.4e-1	В	3.4e-1	В	7.2e0	В	1.2e-7	В	9.5e-7
Arsenic (As)	<,B	2.0e-1	<,B	2.8e-1	<,B	2.8e-1	<,B	5.9e0	<,B	9.9e-8	<,B	7.8e-7
Barium (Ba)	В	4.9e-1	В	6.9e-1	В	7.0e-1	В	1.5e1	В	2.5e-7		1.9e-6
Beryllium (Be)	<,B	1.0e-1	<,B	1.5e-1	<,B	1.5e-1	<,B	3.1e0	<,B	5.2e-8	<,B	4.1e-7
Cadmium (Cd)	В	9.1e-2	В	1.3e-1	В	1.3e-1	В	2.8e0	В	4.6e-8	В	3.6e-7
Chromium (Cr)		2.7e-1		3.8e-1		3.8e-1		8.0e0		1.3e-7		1.1e-6
Cobalt (Co)	В	6.2e-1	В	8.7e-1	В	8.8e-1	В	1.9e1	В	3.1e-7	В	2.5e-6
Copper (Cu)	В	9.9e-1	В	1.4e0	В	1.4e0	В	3.0e1	В	5.0e-7	В	3.9e-6
Lead (Pb)	<,B	2.0e-1	<,B	2.8e-1	<,B	2.8e-1	<,B	6.0e0	<,B	1.0e-7		7.9e-7
Manganese (Mn)		3.9e0		5.6e0		5.7e0		1.2e2		2.0e-6		1.6e-5
Mercury (Hg)		2.3e1		3.2e1		3.3e1		6.9e2		1.1e-5		9.1e-5
Nickel (Ni)	В	3.5e-1	В	5.0e-1	В	5.0e-1	В	1.1e1	В	1.8e-7	В	1.4e-6
Selenium (Se)	В	2.3e-11	В	3.2e-11	В	3.3e-11	В	6.9e-10	В	1.1e-17	В	9.1e-17
Silver (Ag)	<	6.0e-1	<	8.4e-1	<	8.5e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Thallium (TI)	<	5.7e-1	<	8.1e-1	<	8.2e-1	<	1.7e1	<	2.9e-7	<	2.3e-6
Vanadium (V)	<	6.2e-1	<	8.7e-1	<	8.8e-1	<	1.9e1	<	3.1e-7	l	2.5e-6
Zinc (Zn)		1.3e1		1.8e1		1.8e1		3.8e2		6.4e-6		5.1e-5

Table B-14. 0060-END-1.

#### METHOD 0060 SAMPLING DATA SHEET FOR HLLWE TESTS

			IVI	ETHOD 0060	USAM	PLING	DATA	SHEE	I FOR	HLLW	E IES	15				
Site: 1	LLWE Offg	as Tie-in	Sampling	Location:	MA	N-OFG-73	Nozzle No	:		2-01	Est. ΔP:	0.17	Est. Tstac	k, °F:		133
Project:	01-1062	-01-0866	Duct ID, i			12	Nozzle Siz	e, in.:		0.3140	Est. K:	7.66	Est. vs, f	t/s:		27.6
Date:		6/5/2001		ssure, in. WG:			Pitot No.:				Est. ∆H:	1.20	Operator	(s):	FE/F	AL/WS
Run No.:		0-END-1	Est. O2, 9	· · · · · · · · · · · · · · · · · · ·		20.6	Pitot Coeff				Est. DGM 1		e, °F			80
Run Type		Test	Est CO2,			0	Meter Box	No.			Leak Chec					
Pbar., in. Tambient, <sup>c</sup>		25.210 68	Est. Mois Impinger			1.3% 8	∆H@: Y-factor:			1.5673		Pre-		Post-		Dass .
	. Goal (m <sup>3</sup> )	3.0		Goal (ft <sup>3</sup> ):			Min. endin	a DGM vol	(ft <sup>3</sup> )·		DGM Pre: DGM Post:		cfm @ cfm @	1		inHg inHg
Sampling	Clock	Velocity	Meter	Meter	<u> </u>	127.1	<u> </u>	ATURE (°F)	(1. /.	400.700	Pump	0.000	T Citil (a)			illig
Time	Time	ΔΡ΄	ΔН	Volume	Heated			eter		Impinger	Vacuum	%I <sub>i</sub>		COMMEN	ITS	
(min.)	(24hr)	(in. WG)	(in. WG)	(cubic feet)	Line	Stack	1n	Out	Filter	Exit	(in. Hg)					
0	16:00	0.160	1.20	276.678	257	133	69	68	264	48	5.0	-				
10	16:10	0.150	1.20	283,441	259	133	82	69	263	44	5.1	99	O2=20.6			
20	16:20	0.150	1.20	290.220	259	133	. 84	71	262	47	5.1	99	O2=20.5			
30	16:30	0.150	1.20	297.030	259	133	85	72	262	48	5.1	99	<u>L</u> .			
40	16:40	0.150	1.20	303.900	259	133	85	73	262	49	5.1	100	O2=20.5			
50	16:50	0.150	1.20	310.701	259	133	86	73	263	49	5.1	99				
60	17:00	0.150	1.20	317.650	259	133	86	74	263	50	5.1	101	02=20.6			
70	17:10	0.150	1.20	324.560	259	133	. 87	74	263	50	5.1	101				
80	17:20	0.150	1.20	331.653	259	133	87	74	264	51	5.1	103	O2=20.5			
90	17:30	0.150	1.20	338.761	259	133	87	74	263	51	5.1	103				
100	17:40	0.150	1.20	345.873	259	133	86	74	264	52	5.2	104	O2=20.6			
110	17:50	0.150	1.20	352.001	259	133	86	74	263	52	5.2	89				
120	18:00	0.150	1.20	360.078	259	133	86	74	263	52	5.2	118	O2=20.6			
130	18:10	0.150	1.20	367.189	259	133	87	74	261	52	5.2	104				
140	18:20	0.150	1.20	374.275	259	134	87	74	263	53	5.2	103	O2=20.5			
150	18:30	0.150	1.20	381.374	259	134	87	74	264	53	5.2	103				
160	18:40	0.150	1.20	388.481	259	133	87	74	263	53	5.2	103	Q2=20.5			
170	18:50	0.150	1.20	395.590	259	133	86	74	262	53	5.2	104				
180	19:00	0.150	1.20	402.710	259	133	86	74	264	53	5.2	104	O2=20.6			
190	19:10	0.150	1.20	409.800	259	133	86	74	263	53	5.2	103				
200	19:20	0.150	1.20	416.928	259	133	86	74	262	53	5.2	104	O2=20.4			
210	19:30	0.150	1.20	442.102	259	133	86	74	263	53	5.2	367	O2=20.5			
215	19:35	0.150	1.20	427.542	259	133	86	74	263	53	5.2	-424				
Total	Total	ΔPavg	Average	Total		Av	erage Tem	peratures (	°F)		Max.	Ave. %li		-		
215	3:35	0.150	1.20	150.864	259	133	85	73	263	51	5.2	90				

Table B-14. 0060-END-1.

 $O_2\%$ 

CO<sub>2</sub>%

20.5

0.0

## 0060 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site: HLLWE Offgas Tie-in IMPINGER BOX NO. = 8 6/5/2001 Date: Run No.: 0060-END-1 Mercury-Only Section KO-1 Component: Imp-1 Imp-2 Imp-3 Imp-4 Imp-5 Acid Scrub Section Type: short stem modified G-S modified modified short stem modified modified 5% HNO<sub>3</sub> / 10% H<sub>2</sub>O<sub>2</sub> Reagent: None None 4% KMnO<sub>4</sub> / 10% H<sub>2</sub>SO<sub>4</sub> None 2N NaOH Silica Gel Nominal Contents: **Empty** 100 mL solution in each 200 mL solution in each **Empty Empty 100** mL 300-400g Post-test Wt., g: 572.2 717.8 705.1 617.9 732.4 731.7 590.9 678.0 831.3 Impinger Pre-test Wt., g: 572.2 720.5 700.8 617.2 731.7 730.6 590.0 680.3 806.3 wt. gain (g) Wt. Gain, g: 0.0 -2.7 4.3 0.7 0.7 1.1 0.9 -2.3 25.0 27.7 Post-test Volume: 0.0 92.0 102.0 0.0 100.0 100.0 0.0 Impinger Pre-test Volume: 0.0 100.0 100.0 0.0 100.0 100.0 0.0 vol. gain (mL) Volume Gain: 0.0 -8.0 2.0 0.0 0.0 0.0 0.0 -6.0 Post-test pH: 14.0 Filter Lot # 53322 H<sub>2</sub>SO<sub>4</sub> Lot # HNO<sub>3</sub> Lot # 3280601 129100 DI water\* Lot # QCLAB-1 H<sub>2</sub>O<sub>2</sub> Lot # KMnO₄ Lot # 992809 006655 \* used to dilute the other reagents

Table B-14. 0060-END-1.

Project: 01-1062-01-0866										
Run Date:		6/5/2001								
Run Identification:										
PARAMETER	SYMBOL		22.022							
Absolute Pressure in the Duct	Pabs _	in. Hg _	23.923							
Average Duct Gas Temperature	Ts	R	593							
Average Meter Temperature	Tm	R	539							
Average Gas Oxygen Content	Co2,m	%	20.5							
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0							
Total Impinger Weight Gain (water)	Ww	grams	27.7							
Nozzle Area	An	ft²	0.000538							
Duct Area	As	ft²	0.785							
Sample Volume	VmStd	dscf	129.00							
Sample Volume (SI)	VmStdm	dscm	3.653							
Average Sampling Rate	Qm	dscf/m	0.600							
Volume of Water Vapor	VwStd	scf	1.306							
Volume of Water Vapor (SI)	VwStdm	scm	0.0370							
Moisture Fraction	Bws	-	0.010							
Dry Gas Molecular Weight	Md	g/g-mol	28.82							
Wet Gas Molecular Weight	Ms	g/g-mol	28.71							
Gas Velocity at Nozzle	vn	ft/s	25.9							
Gas Velocity at Nozzle (SI)	vnm	m/s	7.89							
Average Gas Velocity	vncor	ft/s	21.84							
Dry Offgas Flow Rate	Qsd	dscf/h	43,505							
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,232							
Actual Offgas Flow Rate	Q	acf/h	61,741							
Intermediate Isokinetic Rate	li	%	102.3							
Final Isokinetic Rate	I	%	102.0							

Table B-14. 0060-END-1.

Project: <sub>01-1062-01-0866</sub>

8/28/2001

Final

6/5/2001 Run Date:

Run Identification: 0060-END-1 Run Type: Test

Lab Report Date: Lab Report Status:

**RESULTS** 

• without blank corrections

(preliminary or final)												
	CONCENTRATIONS				MASS FLOW RATES							
		Actual		Standard	Dry Standard							
		(µg/acm)	(µg/scm)		(µg/dscm)		µg/min		grams/sec		lb/h	
Aluminum (Al)		5.2e1		7.3e1		7.4e1		1.5e3		2.5e-5		2.0e-4
Antimony (Sb)	В	9.8e-1	В	1.4e0	В	1.4e0	В	2.9e1	В	4.8e-7	В	3.8e-6
Arsenic (As)	<	3.1e-1	<	4.3e-1	<	4.4e-1	<	9.0e0	<	1.5e-7	<	1.2e-6
Barium (Ba)	В	1.9e0	В	2.7e0	В	2.7e0	В	5.6e1	В	9.4e-7	В	7.4e-6
Beryllium (Be)	<,B	1.3e-1	<,B	1.8e-1	<,B	3 1.8e-1	<,B	3.7e0	<,B	6.1e-8	<,B	4.8e-7
Cadmium (Cd)	В	1.6e-1	В	2.2e-1	В	2.2e-1	В	4.6e0	В	7.7e-8	В	6.1e-7
Chromium (Cr)		9.1e-1		1.3e0		1.3e0		2.6e1		4.4e-7		3.5e-6
Cobalt (Co)	<	5.2e-1	<	7.3e-1	<	7.4e-1	<	1.5e1	<	2.5e-7	<	2.0e-6
Copper (Cu)	В	6.9e-1	В	9.8e-1	В	9.9e-1	В	2.0e1	В	3.4e-7	В	2.7e-6
Lead (Pb)	В	2.5e-1	В	3.5e-1	В	3.6e-1	В	7.3e0	В	1.2e-7	В	9.7e-7
Manganese (Mn)		4.4e0		6.2e0		6.3e0		1.3e2		2.2e-6		1.7e-5
Mercury (Hg)		2.9e1		4.1e1		4.1e1		8.4 <b>e</b> 2		1.4e-5	ŀ	1.1e-4
Nickel (Ni)	В	1.2e0	В	1.6e0	В	1.6e0	В	3.4e1	В	5.6e-7	В	4.5e-6
Selenium (Se)	<	4.1e-1	<	5.7e-1	<	5.7e-1	<	1.2e1	<	2.0e-7	<	1.6e-6
Silver (Ag)	<	5.0e-1	<	7.0e-1	<	7.1e-1	<	1.5e1	<	2.4e-7	<	1.9e-6
Thallium (TI)	<	4.6e-1	<	6.5e-1	<	6.6e-1	<	1.3e1	<	2.2e-7	<	1.8e-6
Vanadium (V)	<	5.2e-1	<	7.3e-1	<	7.4e-1	<	1.5e1	<	2.5e-7	<	2.0e-6
Zinc (Zn)		2.3e1		3.3e1		3.3e1		6.7e2		1.1e-5		8.9e-5

Table B-14. 0060-END-1.

Project: 01-1062-01-0866

Run Date:

6/5/2001

Run Type:

Run Identification: 0060-END-1 Test

Lab Report Date: Lab Report Status: 8/28/2001

Final

#### **RESULTS**

#### CORRECTED FOR CONTRIBUTION FROM

Corrected for Reagent Blank per EPA SW-846 Test Method 0060 • final presentation should be rounded to two significant digits

(preliminary or final)												
		CC	ONC	ENTRATIO	NS			MA	SS I	FLOW RAT	ES	
		Actual		Standard	Dr	y Standard						
	(	(µg/acm)		(µg/scm)	(	µg/dscm)		μg/min	gı	rams/sec		lb/h
Aluminum (Al)		4.9e1		6.9e1		7.0e1		1.4e3		2.4e-5		1.9e-4
Antimony (Sb)	В	3.0e-1	В	4.2e-1	В	4.2e-1	В	8.7 <b>e</b> 0	В	1.4e-7	В	1.1e-6
Arsenic (As)	<	1.7e-1	<	2.3e-1	<	2.4e-1	<	4.8e0	<	8.1e-8		6.4e-7
Barium (Ba)	В	1.1e0	В	1.5e0	В	1.5e0	В	3.1e1	В	5.2e-7	i	4.1e-6
Beryllium (Be)	<,B	8.7e-2	<,E	3 1.2e-1	<,B	1.2e-1	<,B			4.2e-8	ı	3.3e-7
Cadmium (Cd)	В	1.5e-1	В	2.1e-1	В	2.1e-1	В	4.3e0	В	7.1e-8	ı	5.7e-7
Chromium (Cr)		5.5e-1		7.8e-1		7.9e-1		1.6e1		2.7e-7		2.1e-6
Cobalt (Co)	<	5.2e-1	<	7.3e-1	<	7.4e-1	<	1.5e1		2.5e-7	<	2.0e-6
Copper (Cu)	В	6.9e-1	В	9.8e-1	В	9.9e-1	В	2.0e1	В	3.4e-7	ı	2.7e-6
Lead (Pb)	В	1.7e-1	В	2.4e-1	В	2.4e-1	В	4.9e0	В	8.2e-8	В	6.5e-7
Manganese (Mn)		4.2e0		5.9e0		6.0e0		1.2e2		2.0e-6		1.6e-5
Mercury (Hg)		2.9e1		4.1e1		4.1e1		8.4e2		1.4e-5		1.1e-4
Nickel (Ni)	В	4.9e-1	В	6.9e <b>-1</b>	В	7.0e-1	В	1.4e1	В	2.4e-7	В	1.9e-6
Selenium (Se)	<	1.9e-11	<	2.7e-11	<	2.7e-11	<	5.6e-10	<	9.4e-18	<	7.4e-17
Silver (Ag)	<	5.0e-1	<	7.1e-1	<	7.1e-1	<	1.5e1	<	2.4e-7	<	1.9e-6
Thallium (TI)	<	4.6e-1	<	6.5e-1	<	6.6e-1	<	1.3e1	<	2.2e-7	<	1.8e-6
Vanadium (V)	<	5.2e-1	<	7.3e-1	<	7.4e-1	<	1.5e1	<	2.5e-7	<	2.0e-6
Zinc (Zn)		2.2e1		3.1e1		3.1e1		6.4e2		1.1e-5		8.5e-5

Table B-15. 0060-STRT-2.

#### METHOD 0060 SAMPLING DATA SHEET FOR HLLWE TESTS

			1911	ETHOD UUGU	JAIVII	LING	DAIA	JIILL	1101	IILLVV	LILU				
Site:	HLLWE C	Offgas Tie-in	Sampling	Location:	MA	N-OFG-73	Nozzle No	.:			Est. ΔP:	0.16	Est. Tstack, °F:		133
Project:	01-10	062-01-0866	Duct ID, it	nches:		12	Nozzle Siz	e, in.:		0.3140		7.67	Est. vs, ft/s:		26.7
Date:				ssure, in. WG:		-17.5	Pitot No.:				Est. ∆H:		Operator(s):	RV	/,FE, JA
Run No.:		060-STRT-2		-		20.6	Pitot Coeff				Est. DGM 1		e, °F		80
Run Type		Test	Est CO <sub>2</sub> ,			0	Meter Box	No.			Leak Chec				
Pbar., in. I		25.219	Est. Moist	<del></del>			ΔH@:			1.5673		Pre-	PASS Post-		PASS
Tambient, °	F: Goal (m <sup>3</sup> ):	3.0	Impinger	Box No.: Goal (ft³):		9 127.1	Y-factor:	g DGM vol.	/ft <sup>3</sup> \·		DGM Pre: DGM Post:		cfm @	17 6	inHg inHg
			<u> </u>			127.1			π. (ιτ ).	334.002		0.000	T		iiiiig
Sampling Time	Clock Time	Velocity ΔP	Meter ΔH	Meter Volume	Heated			ATURE (°F)		Impinger	Pump Vacuum	%l <sub>i</sub>	COM	MENTS	
(min.)	(24hr)	(in. WG)	(in. WG)	(cubic feet)	Line	Stack	In	Out	Filter	Exit	(in. Hg)				
0	7:51	0.170	1.20	427.722	255	134	63	53	260	48	5.3				
10	8:01	0.170	1.30	434.370	261	134	74	58	261	45	5.5	94			
20	8:11	0.170	1.30	441.310	260	134	76	62	265	49	5.5	97			
30	8:21	0.170	1.30	448.280	260	134	77	63	264	49	5.5	97			
40	8:31	0.170	1.30	455.250	260	134	78	64	261	50	5.5	97			
50	8:41	0.170	1.30	462.210	258	134	81	67	261	51	5.5	96			
59	8:50	0.170	1.30	468.605	258	134	82	68	263	52	5.5	98	Stopped train beca	ause of	NWCF
59	9:22	0.170	1.30	468.605	255	134	78	67	264	53	5.5	#DIV/0!	building evacuation	n.	
60	9:23	0.170	1.30	469.290	255	134	79	67	264	53	5.5	95_	Restart at 0922		
70	9:33	0.170	1.30	476.420	250	134	84	68	262	49	5.5	98	Hood O2=20.5		
80	9:43	0.170	1.30	483.530	252	134	85	69	263	52	5.5	98	Hood O2=20.4		
90	9:53	0.170	1.30	490.650	257	134	85	70	264	53	5.5	98	Hood O2=20.4		
100	10:03	0.170	1.30	497.800	254	134	86	71	262	53	5.5	98	Hood O2=20.5		
110	10:13	0.170	1.30	504.940	255	134	86	72	263	54	5.5	98	Offgas O2=20.5		
120	10:23	0.170	1.30	512.090	251	134	87	73	262	54	5.5	98	Offgas O2=20.5		
130	10:33	0.160	1.30	519.430	250	133	88	73	262	55	5.5	103			
140	10:43	0.160	1.30	527.621	252	133	88	74	262	54	5.5	115			
150	10:53	0.160	1.30	533,730	250	134	88	73	261	55	5.5	86			
160	11:03	0.160	1.30	540.740	254	134	88	75	261	56	5.5	99	Offgas O2=20.5		
170	11:13	0.160	1.25	548.003	251	133	89	76	263	56	5.5	102	Offgas O2≃20.5		
180	11:23	0.160	1.25	555.122	253	133	89	76	261	56	5.5	100	Offgas O2=20.5		
190	11:33	0.160	1.30	562.328	250	133	90	77	262	57	5.5	101			
Total	Total	ΔPavg	Average	Total	1	A	verage Tem	peratures (	(°F)	1	Max.	Ave. %l;	<del> </del>		
190	3:10	0.167	1.29	134.606	255	134	83	69	262	52	5.5	#DIV/0!			

Table B-15. 0060-STRT-2.

# 0060 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site: HLLWE Offgas Tie-in IMPINGER BOX NO. = Date: 6/6/2001 0060-STRT-2 Run No.: Mercury-Only Section Acid Scrub Section Component: KO-1 lmp-1 Imp-2 Imp-3 Imp-4 Imp-5 modified G-S modified modified modified Type: short stem modified short stem Reagent: None 5% HNO<sub>3</sub> / 10% H<sub>2</sub>O<sub>2</sub> None 4% KMnO<sub>4</sub> / 10% H<sub>2</sub>SO<sub>4</sub> None 2N NaOH Silica Gel 300-400g **Nominal Contents:** 100 mL solution in each 100 mL solution in each 100 mL **Empty Empty** Empty Post-test Wt., g: 559.6 723.0 678.4 583.4 739.1 732.5 567.0 709.6 813.0 Impinger Pre-test Wt., g: 559.6 726.2 674.0 582.4 739.6 733.3 564.3 716.1 785.1 wt. gain (g) Wt. Gain, g: 0.0 -3.2 4.4 1.0 -0.5 -0.8 2.7 -6.5 27.9 25.0 Post-test Volume: 0.0 100.0 100.0 0.0 100.0 100.0 0.0 Impinger Pre-test Volume: 0.0 100.0 100.0 0.0 100.0 100.0 0.0 vol. gain (mL) Volume Gain: 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 14.0 Post-test pH: H<sub>2</sub>SO<sub>4</sub> Lot # Filter Lot # 53322 328060 HNO<sub>3</sub> Lot # 129100 DI water\* Lot # QCLAB-1 H<sub>2</sub>O<sub>2</sub> Lot # KMnO<sub>4</sub> Lot # 992809 006655

\* used to dilute the other reagents

 $\frac{O_2\%}{CO_2\%}$  20.5 0.0

Table B-15. 0060-STRT-2.

## FIELD DATA CALCULATIONS

Project: 01-1062-01-0866 Run Date: 6/6/2001											
Run Date: Run Identification:		200 T STRT-2									
PARAMETER	SYMBOL	UNITS									
Absolute Pressure in the Duct	Pabs	in. Hg	23.932								
Average Duct Gas Temperature	Ts	R	593								
Average Meter Temperature	Tm	R	536								
Average Gas Oxygen Content	Co2,m	%	20.5								
Average Gas Carbon Dioxide Content	Cco2,m	%	0.0								
Total Impinger Weight Gain (water)	Ww	grams	25.0								
Nozzie Area	An	ft²	0.000538								
Duct Area	As	ft²	0.785								
Sample Volume	VmStd	dscf	115.90								
Sample Volume (SI)	VmStdm	dscm	3.282								
Average Sampling Rate	Qm	dscf/m	0.610								
Volume of Water Vapor	VwStd	scf	1.179								
Volume of Water Vapor (SI)	VwStdm	scm	0.0334								
Moisture Fraction	Bws	-	0.010								
Dry Gas Molecular Weight	Md	g/g-mol	28.82								
Wet Gas Molecular Weight	Ms	g/g-mol	28.71								
Gas Velocity at Nozzle	vn	ft/s	27.3								
Gas Velocity at Nozzle (SI)	vnm	m/s	8.31								
Average Gas Velocity	vncor	ft/s	23.00								
Dry Offgas Flow Rate	Qsd	dscf/h	45,790								
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,297								
Actual Offgas Flow Rate	Q	acf/h	65,037								
Intermediate Isokinetic Rate	li	%	98.8								
Final Isokinetic Rate	1	%	98.5								

Table B-15. 0060-STRT-2.

Project: 01-1062-01-0866

Run Date:

6/6/2001

Run Type: Test

Run Type: Lab Report Date: Test 8/28/2001

Lab Report Status: (preliminary or final)

Final

#### **RESULTS**

#### • without blank corrections

• final presentation should be rounded to two significant digits

(preminary or miar)												~~
		CC	NC	ENTRATIO	NS			M.A	SS	FLOW RAT	ES	
		Actual	S	Standard	Dry	Standard	l					
	(1	µg/acm)	(	(µg/scm)	(þ	ıg/dscm)		μg/min	gı	rams/sec		lb/h
Aluminum (Al)		1.6e1		2.3e1		2.3e1		5.0e2		8.3e-6		6.6e-5
Antimony (Sb)	В	9. <b>7</b> e-1	В	1.4e0	В	1.4e0	В	3.0e1	В	4.9e-7	В	3.9e-6
Arsenic (As)	<	3.4e-1	<	4.8e-1	<	4.9e-1	<	1.1e1	<	1.8e-7	<	1.4e-6
Barium (Ba)	В	1.1e0	В	1.5e0	В	1.6e0	В	3.4e1	В	5.6e-7	В	4.4e-6
Beryllium (Be)	<,B	1.4e-1	<,B	2.0e-1	<,B	2.0e-1	<,B	4.3e0	<,B	7.1e-8	<,B	5.7e-7
Cadmium (Cd)	<,B	5.8e-2	<,B	8.1e <b>-</b> 2	<,B	8.2e-2	<,B	1.8e0	<,B	3.0e-8	<,B	2.4e-7
Chromium (Cr)		5.4e-1		7.5e-1		7.6e-1		1.6e1		2.7e-7	•	2.2e-6
Cobalt (Co)	<	5.8e-1	<	8.1e-1	<	8.2e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Copper (Cu)	В	4.9e-1	В	6.9e-1	В	7.0e-1	В	1.5e1	В	2.5e-7	В	2.0e-6
Lead (Pb)	<,B	3.0e-1	<,B	4.2e-1	<,B	4.3e-1	<,B	9.2e0	<,B	1.5e-7	<,B	1.2e-6
Manganese (Mn)		8.2e0		1.1e1		1.2e1		2.5e2		4.2e-6		3.3e-5
Mercury (Hg)		2.4e1		3.3e1		3.4e1		7.2e2		1.2e-5		9.6e-5
Nickel (Ni)	В	9.7e-1	В	1.4e0	В	1.4e0	В	3.0e1	В	4.9e-7	В	3.9e-6
Selenium (Se)	В	5.6e-1	В	7.8e-1	В	7.9e-1	В	1.7e1	В	2.9e-7	В	2.3e-6
Silver (Ag)	<	5.6e-1	<	7.8e-1	<	7.9e-1	<	1.7e1	<	2.9e-7	<	2.3e-6
Thallium (TI)	<	5.4e-1	<	7.5e-1	<	7.6e-1	<	1.6e1	<	2.7e-7	<	2.2e-6
Vanadium (V)	<	5.8e-1	<	8.1e-1	<	8.2e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Zinc (Zn)		6.4e0		9.0e0		9.1e0		2.0e2		3.3e-6		2.6e-5

Table B-15. 0060-STRT-2.

Project: 01-1062-01-0866

Run Date: 6/6/2001 Run Identification: 0060-STRT-2

Run Type: Lab Report Date: Lab Report Status:

Test 8/28/2001

Final

#### **RESULTS**

## CORRECTED FOR CONTRIBUTION FROM

Corrected for Reagent Blank per EPA SW-846 Test Method 0060 • final presentation should be rounded to two significant digits

(preliminary or final)		Final										
(preliminary or imal)	I	CC	NCI	ENTRATIO	NS			MA	SS F	LOW RAT	ES	
	<b> </b> -	Actual		Standard		Standard						
	l	ug/acm)		µg/scm)		ıg/dscm)	l.	µg/min	gr	ams/sec		lb/h
Aluminum (Al)		1.5e1		2.1e1		2.1e1		4.5e2		7.5e-6		6.0e-5
Antimony (Sb)	В	2.0e-1	В	2.8e-1	В	2.9e-1	В	6.2e0	В	1.0e-7	В	8.2e-7
Arsenic (As)	<	1.8e-1	<	2.6e-1	<	2.6e-1	<	5. <b>7e</b> 0	<	9.4e-8		7.5e-7
Barium (Ba)	В	1.2e-1	В	1.6e-1	В	1.6e-1	В	3.6e0	В	5.9e-8	В	4.7e-7
Beryllium (Be)	<,B	9.7e-2	<,B	1.4e-1	<,B	1.4e-1	<,B	3.0e0	<,B	4.9e-8	<,B	3.9e-7
Cadmium (Cd)	<,B	4.5e-2	<,B	6.4e-2	<,B	6.4e-2	<,B	1.4e0	<,B	2.3e-8	<,B	1.8e-7
Chromium (Cr)	-	1.4e-1		2.0e-1		2.0e-1		4.4e0	]	7.4e-8		5.8e-7
Cobalt (Co)	<	5.8e-1	<	8.1e-1	<	8.2e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Copper (Cu)	В	4.8e-1	В	6.8e-1	В	6.9e-1	В	1.5e1	В	2.5e-7	В	2.0e-6
Lead (Pb)	<,B	2.1e-1	<,B	3.0e-1	<,B	3.0e-1	<,B	6.5e0	<,B	1.1e-7	<,B	8.5e-7
Manganese (Mn)		7.8e0		1.1e1		1.1e1		2.4e2		4.0e-6		3.2e-5
Mercury (Hg)		2.4e1		3.3e1		3.4e1		7.2e2		1.2e-5		9.6e-5
Nickel (Ni)	В	2.2e-1	В	3.1e-1	В	3.2e-1	В	6.8e0	В	1.1e-7	В	9.1e-7
Selenium (Se)	В	2.1e-11	В	3.0e-11	В	3.0e-11	В	6.6e-10	В	1.1e-17	В	8.7e-17
Silver (Ag)	<	5.6e-1		7.9e-1	<	8.0e-1	<	1.7e1	<	2.9e-7	<	2.3e-6
Thallium (TI)	<	5.4e-1	<	7.5e-1	<	7.6e-1	<	1.6e1	<	2.7e-7	<	2.2e-6
Vanadium (V)	<	5.8e-1	<	8.1e-1	<	8.2e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Zinc (Zn)		5.6e0		7.8e0		7.9e0		1.7e2		2.8e-6		2.3e-5

		·

Table B-16. 0060-END-2.

#### METHOD 0060 SAMPLING DATA SHEET FOR HILLWE TESTS

	METHOD 0060 SAMPLING DATA SHEET FOR HLLWE TESTS														
Site: I	LLWE Offg	as Tie-in	Sampling	Location:	MAI	V-OFG-73	Nozzle No	.:			Est. ∆P:	0.15	Est. Tsta	ock, °F:	133
Project:	-	-01-0866	Duct ID, i	<del></del>		12	Nozzle Siz	e, in.:			Est. K:		Est. vs,		26.7
Date:		6/6/2001		essure, in. WG:		-17.5	Pitot No.:				Est. ∆H:		Operato	or(s):	FE/RW
Run No.:		0-END-2	Est. O <sub>2</sub> , 9			20.5	Pitot Coeff		a.		Est. DGM		re, °F		80
Run Typ		Test	Est CO <sub>2</sub> ,			0	Meter Box	No.			Leak Chec				
Pbar., in.	_ <del>_</del>	25.192	Est. Mois				ΔH@:			1.5673		Pre-	Pass	Post-	Pass
Tambient, <sup>o</sup>	. Goal (m³)	70 3.0	Impinger DGM vol	Goal (ft'):		8 127.1	Y-factor: Min. endin	a DGM vo	l (fft)·		DGM Pre: DGM Post:		cfm @ cfm @	15.5	inHg
						127.1				090.100	L	0.000	Cilli @	6	inHg
Sampling Time	Clock Time	Velocity ΔP	Meter ∆H	Meter Volume	Heated			TURE (°F)		Impinger	Pump Vacuum	%l,		COMMENTS	ļ
(min.)	(24hr)	(in. WG)	(in. WG)	(cubic feet)	Line	Stack	In	Out	Filter	Exit	(in. Hg)	<u> </u>			
0	15:00	0.150	1.25	563.026	248	133	82	75	257	57	5.2				
10	15:10	0.150	1.25	569.885	248	133	87	76	259	55	5.2	100			
20	15:20	0.150	1.25	576.880	250	133	90	78	260	55	5.2	101	O2=20.5		
30	15:30	0.150	1.25	583.890	249	133	91	79	260	55	5.2	101	O2=20.5		
40	15:40	0.150	1.20	590.900	251	133	92	79	259	55	5.2	101	O2=20.5		
50	15:50	0.150	1.20	597.920	250	133	93	80	261	55	5.2	101	O2=20.5		
60	16:00	0.150	1.20	604.930	250	133	93	80	261	55	5.2	101_	O2=20.5		
70	16:10	0.150	1.20	611.920	254	133	93	81	260	56	5.2	100	O2=20.4		
80	16:20	0.150	1.20	618.940	254	133	93	81	260	57	5.2	101	O2=20.5		
90	16:30	0.150	1.20	625.950	256	133	93	81	260	57	5.2	101	O2=20.4		
100	16:40	0.150	1.20	632.960	256	133	93	81	260	58	5.2	101	02=20.5		
110	16:50	0.150	1.20	639.960	256	133	93	81	260	58	5.2	101	02=20.4		
120	17:00	0.150	1.20	646.980	255	133	93	81	261	59	5.2	101	O2=20.4		
130	17:10	0.150	1.20	654.010	256	133	93	81	261	59	5.2	101	O2=20.4		
140	17:20	0.150	1.20	661.020	255	133	93	81	259	60	5.2	101	02=20.4		
150	17:30	0.150	1.20	668.060	255	133	93	81	260	60	5.2	101	O2=20.5		
160	17:40	0.150	1.20	675.060	255	133	93	81	260	61	5.2	101	O2=20.4		
170	17:50	0.150	1.20	682.070	255	133	93	81	260	61	5.2	101	O2=20.4		
180	18:00	0.150	1.30	689.520	255	133	93	81	260	61	5.5	107	O2=20.5		
182	18:02	0.150	1.30	691.050	255	133	93	81	260	61	5.5	110	02=20.4		
Total	Total	ΔPavg	Average	Total		A	verage Tem	peratures	(°F)		Max.	Ave. %l <sub>i</sub>	<u> </u>		
182	3:02	0.150	1.22	128.024	253	133	92	80	260	58	5.5	102			

Table B-16. 0060-END-2.

# 0060 CONFIGURATION TRAIN COMPONENT DATA SHEET for HLLWE OFFGAS SAMPLING

Site: HLLWE Offgas Tie-in IMPINGER BOX NO. = 8 6/6/2001 Date: 0060-END-2 Run No.: Mercury-Only Section Acid Scrub Section Component: KO-1 Imp-1 Imp-2 Imp-3 Imp-4 lmp-5 modified modified modified modified G-S modified short stem Type: short stem 5% HNO<sub>3</sub> / 10% H<sub>2</sub>O<sub>2</sub> None 4% KMnO<sub>4</sub> / 10% H<sub>2</sub>SO<sub>4</sub> None 2N NaOH Silica Gel Reagent: None 100 mL 300-400g 100 mL solution in each 200 mL solution in each **Empty** Nominal Contents: Empty **Empty** 734.6 718.7 608.8 691.6 781.0 Impinger Post-test Wt., g: 572.2 720.7 682.1 618.4 Pre-test Wt., g: 573.2 727.9 677.4 618.6 734.9 719.0 607.3 694.7 754.2 wt. gain (g) 20.9 Wt. Gain, g: -1.0 -7.2 4.7 -0.2 -0.3 -0.3 1.5 -3.1 26.8 Post-test Volume: 0.0 100.0 100.0 0.0 100.0 100.0 0.0 Impinger 0.0 100.0 100.0 0.0 100.0 100.0 0.0 vol. gain (mL) Pre-test Volume: Volume Gain: 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 13.0 Post-test pH: H<sub>2</sub>SO<sub>4</sub> Lot # 328060 Filter Lot # 53322 HNO<sub>3</sub> Lot # 129100 DI water\* Lot # QCLAB1 KMnO₄ Lot # H<sub>2</sub>O<sub>2</sub> Lot # 992809 06655

<sup>\*</sup> used to dilute the other reagents

$O_2\%$	20.5
CO <sub>2</sub> %	0.0

Table B-16. 0060-END-2.

## FIELD DATA CALCULATIONS

Project:	01-1062		
Run Date:			
Run Identification: PARAMETER	SYMBOL	END-2 UNITS	<u> </u>
Absolute Pressure in the Duct	Pabs	in. Hg	23.905
Average Duct Gas Temperature	Ts	R	593
Average Meter Temperature	Tm	R	546
Average Gas Oxygen Content	Co2,m	%	20.5
	·	% %	0.0
Average Gas Carbon Dioxide Content	Cco2,m		
Total Impinger Weight Gain (water)	Ww	grams	20.9
Nozzle Area	An	ft²	0.000538
Duct Area	As	ft²	0.785
Sample Volume	VmStd	dscf	108.05
Sample Volume (SI)	VmStdm	dscm	3.060
Average Sampling Rate	Qm	dscf/m	0.594
Volume of Water Vapor	VwStd	scf	0.985
Volume of Water Vapor (SI)	VwStdm	scm	0.0279
Moisture Fraction	Bws	-	0.009
Dry Gas Molecular Weight	Md	g/g-mol	28.82
Wet Gas Molecular Weight	Ms	g/g-mol	28.72
Gas Velocity at Nozzle	vn	ft/s	25.8
Gas Velocity at Nozzle (SI)	vnm	m/s	7.88
Average Gas Velocity	vncor	ft/s	21.81
Dry Offgas Flow Rate	Qsd	dscf/h	43,465
Dry Offgas Flow Rate (SI)	Qsdm	dscm/h	1,231
Actual Offgas Flow Rate	Q	acf/h	61,660
Intermediate Isokinetic Rate	li	%	101.4
Final Isokinetic Rate	l	%	101.0

Table B-16. 0060-END-2.

Project: 01-1062-01-0866

Run Date:

6/6/2001

Run Identification: 0060-END-2 Run Type:

Test 8/28/2001

Lab Report Date: Lab Report Status: (preliminary or final)

Final

#### **RESULTS**

#### • without blank corrections

• final presentation should be rounded to two significant digits

(preliminary or imar)															
		CC	DNC	ENTRATIO	NS			MA	SS	FLOW RAT	ES				
		Actual	,	Standard	Dry	Standard	l								
		(µg/acm)		(µg/scm)	(١	ıg/dscm)		μg/min	gi	rams/sec		lb/h			
Aluminum (AI)		1.7e1		2.4e1		2.4e1		4.9e2		8.2e-6		6.5e-5			
Antimony (Sb)	В	9.4e-1	В	1.3e0	В	1.3e0	В	2.7e1	В	4.6e-7	В	3.6e-6			
Arsenic (As)	<	3.7e-1	<	5.2e-1	<	5.2e-1	<	1.1e1	<	1.8e-7	<	1.4e-6			
Barium (Ba)	В	1.1e0	В	1.5e0	В	1.5e0	В	3.2e1	В	5.3e-7	В	4.2e-6			
Beryllium (Be)	<,B	1.5e-1	<,B	2.1e-1	<,B	2.1e-1	<,B	4.4e0	<,B	7.3e-8	<,B	5.8e-7			
Cadmium (Cd)	<	6.2e-2	<	8.7e-2	<	8.8e <b>-</b> 2	<	1.8e0	<	3.0e-8	<	2.4e-7			
Chromium (Cr)		1.2e0		1.7e0		1.7e0		3.4e1		5.7e-7		4.5e-6			
Cobalt (Co)	<	6.2e-1	<	8.7e-1	<	8.8e-1	<	1.8e1		3.0e-7	<	2.4e-6			
Copper (Cu)	<,B	3.2e-1	<,B	4.5e-1	<,B	4.6e-1	<,B	9.4e0	<,B	1.6e-7	<,B	1.2e-6			
Lead (Pb)	<,B	2.8e-1	<,B	3.9e-1	<,B	3.9e-1	<,B	8.0e0	<,B	1.3e-7	<,B	1.1e-6			
Manganese (Mn)		1.7e1		2.3e1		2.4e1		4.8e2		8.0e-6		6.4e-5			
Mercury (Hg)		2.5e1		3.6e1		3.6e1		7.4e2		1.2e-5		9.8e-5			
Nickel (Ni)	В	9.4e-1	В	1.3e0	В	1.3e0	В	2.7e1	В	4.6e-7	В	3.6e-6			
Selenium (Se)	В	5.3e-1	В	7.4e-1	В	7.5e-1	В	1.5e1	В	2.6e-7	В	2.0e-6			
Silver (Ag)	<	6.0e-1	<	8.4e-1	<	8.5e-1	<	1.7e1	<	2.9e-7	<	2.3e-6			
Thallium (TI)	<	5.5e-1	<	7.8e-1	<	7.8e-1	<	1.6e1	<	2.7e-7	<	2.1e-6			
Vanadium (V)	<	6.2e-1	<	8.7e-1	<	8.8e-1	<	1.8e1	<	3.0e-7	<	2.4e-6			
Zinc (Zn)		3.9e0		5.5e0		5.6e0		1.1e2		1.9e-6		1.5e-5			

Table B-16. 0060-END-2.

Project: <sub>01-1062-01-0866</sub>

Run Date: 6/6/2001

Run Identification: 0060-END-2 Run Type: Test

Lab Report Date: 8/28/2001 Lab Report Status:

(preliminary or final)

# RESULTS CORRECTED FOR CONTRIBUTION FROM

Corrected for Reagent Blank per EPA SW-846 Test Method 0060 • final presentation should be rounded to two significant digits

(preliminary or linar	<i>)</i>			_								
		CC	DNC	<b>ENTRATIO</b>	NS			MA	SS F	LOW RAT	ES	
		Actual	3	Standard	Dry	/ Standard						
	(µ	g/acm)	(	(µg/scm)	(١	ug/dscm)		µg/min	gr	ams/sec		lb/h
Aluminum (Al)		1.5e1		2.1e1		2.2e1		4.4e2		7.4e-6		5.9e-5
Antimony (Sb)	В	1.2e-1	В	1.7e-1	В	1.8e-1	В	3.6e0	В	6.0e-8	В	4.8e-7
Arsenic (As)	<	2.0e-1	<	2.8e-1	<	2.8e-1	<	5.8e0	<	9.6e-8	<	7.6e-7
Barium (Ba)	В	3.2e-2	В	4.5e-2	В	4.6e-2	В	9. <b>4e-1</b>	В	1.6e-8	В	1.2e-7
Beryllium (Be)	<,B	1.0e-1	<,B	1.5e-1	<,B	1.5e-1	<,B	3.0e0	<,B	5.0e-8	<,B	4.0e-7
Cadmium (Cd)	<	4.9e-2	<	6.8e-2	<	6.9e <b>-</b> 2	<	1.4e0	<	2.4e-8	<	1.9e-7
Chromium (Cr)		7.5e-1		1.1e0		1.1e0		2.2e1		3.7e-7		2.9e-6
Cobalt (Co)	<	6.2e-1	<	8.7e-1	<	8.8e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Copper (Cu)	<,B	3.2e-1	<,B	4.4e-1	<,B	4.5e-1	<,B	9.2e0	<,B	1.5e-7	<,B	1.2e-6
Lead (Pb)	<,B	1.8e-1	<,B	2.5e-1	<,B	2.5e-1	<,B	5.2e0	<,B	8.6e-8	<,B	6.8e-7
Manganese (Mn)		1.6e1		2.2e1		2.3e1		4.6e2		7.7e-6		6.1e-5
Mercury (Hg)		2.5e1		3.6e1		3.6e1		7.4e2		1.2e-5		9.8e-5
Nickel (Ni)	В	1.5e-1	В	2.1e-1	В	2.1e-1	В	4.3e0	В	7.2e-8	В	5.7e-7
Selenium (Se)	В	2.3e-11	В	3.2e-11	В	3.3e-11	В	6.7e-10	В	1.1e-17	В	8.9e-17
Silver (Ag)	<	6.0e-1	<	8.5e-1	<	8.5e-1	<	1.7e1	<	2.9e-7	<	2.3e-6
Thallium (TI)	<	5.5e-1	<	7.8e-1	<	7.8e-1	<	1.6e1	<	2.7e-7	<	2.1e-6
Vanadium (V)	<	6.2e-1	<	8.7e-1	<	8.8e-1	<	1.8e1	<	3.0e-7	<	2.4e-6
Zinc (Zn)		2.9e0		4.0e0		4.1e0		8.4e1		1.4e-6		1.1e-5

Table B-17. SVOC emission rates - grames per second comparisons.

Table B-17. SVOC emission rates	s - grames pe	er secona c	compans				I Project			L Project			I Brojost	<del>,</del>		- <del></del>	I Urningt	<u> </u>	EKNI VENO
Analyte	Registry	STRT-1	Flag	Specific	END-1	Flag	Specific	STRT-2	Flag	Specific	END-2	Flag	Specific	Max value ፲ g/s ଓ	Avg+2σ ਜ਼ੁ a/s a	Results a/s	Project Specific	Run Avgs	END Run Avgs
1	Number	g/s	Ğ	Flag	g/s	ĝ	Flag	g/s	ĝ	Flag	g/s	Õ	Flag	g/s 🖔	g/s ຜັ	g/s d	Flag	g/s	g/s
Acenaphthene	83-32-9	1.1e-6	<	N	1.1e-6	<	N	1.8e-6	<,J	P	9.9e-7	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	P	1.4e-6	1.0e-6
Acenaphthylene	208-96-8	1.0e-6	<	l N	1.0e-6	<	N	1.8e-6	<,J	Р	9.6e-7	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	Р	1.4e-6	1.0e-6
Acetophenone	98-86-2	5.4e-6	<,J	Р	3.9e-6	<,J	P	4.8e-6	<,J	Р	3.8e-6	<,J	P	5.4e-6 <,J	6.0e-6	5.4e-6	P	5.1e-6	3.8e-6
Aniline	62-53-3	1.2e-5	<	N	1.2e-5	<	N	1.7e-5	<	N	1.1e-5	<	N	1.7e-5 <	1.8e-5	1.7e-5	N	1.5e-5	1.2e-5
Anthracene	120-12-7	1.0e-6	<	N	1.0e-6	<	N	1.8e-6	<,J	Р	9.6e-7	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	Р	1.4e-6	1.0e-6
Benzidine	92-87-5	7.3e-5	<	N	7.2e-5	<	N	9.1e-5	<	N	6.8e-5	<	N	9.1e-5 <	9.7e-5	9.1e-5	N	8.2e-5	7.0e-5
Benzoic acid	65-85-0	5.4e-4	Ε	P	2.8e-4	Е	Р	2.6e-4	E	P	2.7e-4	E	P	5.4e-4 E	6.0e-4	5.4e-4	Р	4.0e-4	2.8e-4
Benzo(a)anthracene	56-55-3	1.3e-6	<	N	1.3e-6	<	N	2.2e-6	<,J	Р	1.2e-6	<	N	2.2e-6 <,J	2.4e-6	2.2e-6	Р	1.7e-6	1.3e-6
Benzo(a)pyrene	50-32-8	1.4e-5	<	N	1.4e-5	<	N	1.6e-5	<,J	Р	1.3e-5	<	N	1.6e-5 <,J	1.7e-5	1.6e-5	Р	1.5e-5	1.4e-5
Benzo(b)fluoranthene	205-99-2	3.3e-5	<	N	3.2e-5	<	N	3.4e-5	<,J	Р	3.0e-5	<	N	3.4e-5 <,J	3.6e-5	3.4e-5	Р	3.3e-5	3.1e-5
Benzo(g,h,i)perylene	191-24-2	1.9e-5	<	N I	1.8e-5	<	N	2.0e-5	ر,>	P	1.7e-5	<	N	2.0e-5 <,J	2.1e-5	2.0e-5	Р	2.0e-5	1.8e-5
Benzo(k)fluoranthene	207-08-9	4.7e-5	<	N	4.6e-5	<	N	4.9e-5	<,J	P	4.4e-5	<	N	4.9e-5 <,J	5.1e-5	4.9e-5	P	4.8e-5	4.5e-5
Benzyl alcohol	100-51-6	6.2e-5	<	N I	6.2e-5	<	N I	6.5e-5	<	N	5.8e-5	<	N	6.5e-5 <	6.7e-5	6.5e-5	N	6.4e-5	6.0e-5
bis(2-Chloroethoxy)methane	111-91-1	1.1e-6	<	N	1.1e-6	< <	N	1.6e-6	<	N P	1.0e-6	<	N	1.6e-6 <	1.7e-6	1.6e-6	N P	1.3e-6	1.0e-6
bis(2-Chloroethyl)ether bis(2-Ethylhexyl)phthalate	111-44-4 117-81-7	1.2e-6 1.3e-5	<,J	A	1.2e-6 2.4e-5	<	N p	1.8e-6 1.5e-5	<,J		1.1e-6	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	P	1.5e-6	1.2e-6
4-Bromophenyl-phenylether	101-55-3	1.0e-6		N	1.0e-6		N	2.0e-6	ر,> ل,>	A	1.6e-5 9.5e-7	<,J <	A N	2.4e-5 < 2.0e-6 <,J	2.6e-5 2.3e-6	2.4e-5 2.0e-6	P	1.4e-5 1.5e-6	2.0e-5 9.8e-7
Butylbenzylphthalate	85-68-7	1.4e-6	<	N	1.4e-6	<	N	2.0e-6 2.2e-6	<,J	P	1.3e-6	<	N	2.2e-6 <,J	2.4e-6	2.0e-6 2.2e-6		1.8e-6	1.4e-6
Carbazole	86-74-8	1.4e-6	<	N	1.4e-6	<	N	2.2e-6 2.0e-6	<,J	P	1.3e-6	<	N	2.0e-6 <,J	2.2e-6	2.0e-6	P	1.7e-6	1.4e-6
4-Chloro-3-methylphenol	59-50-7	1.9e-6	<	N N	1.8e-6	<	N	3.5e-6	۰,۵ ۲,۶	N	1.7e-6	<	N	3.5e-6 <,J	4.0e-6	3.5e-6	N	2.7e-6	1.8e-6
4-Chloroaniline	106-47-8	9.8e-6	<	N	9.7e-6	~	N	1.3e-5	< .,5	N	9.1e-6	~	N	1.3e-5 <	1.3e-5	1.3e-5	l N	1.1e-5	9.4e-6
2-Chloronaphthalene	91-58-7	1.0e-6	<	N	1.0e-6	<	N	1.7e-6	<,J	"	9.4e-7	~	N	1.7e-6 <,J	1.9e-6	1.7e-6	P	1.1e-5 1.4e-6	9.7e-7
2-Chlorophenol	95-57-8	1.2e-6	<	N	1.2e-6	<	N	1.7e-6	ا,. ال,>	P	1.1e-6	<	l 'n l	1.7e-6 <,J	1.8e-6	1.7e-6	P	1.5e-6	1.2e-6
4-Chlorophenyl phenyl ether	7005-72-36	1.2e-6	<	N	1.2e-6	<	N	1.9e-6	ر. ال,>	P	1.1e-6	<	N	1.9e-6 <,J	2.1e-6	1.9e-6	P	1.6e-6	1.2e-6
Chrysene	218-01-9	1.3e-6	<	N	1.3e-6	<	N	2.4e-6	<,J	Р	1.2e-6	<	N	2.4e-6 <,J	2.7e-6	2.4e-6	P	1.9e-6	1.3e-6
Di-n-butylphthalate	84-74-2	1.1e-5	<,J	P	1.1e-5	<,J	Р	1.3e-5	<,J	P	1.0e-5	<,J	P	1.3e-5 <,J	1.3e-5	1.3e-5	P	1.2e-5	1.0e-5
Di-n-octylphthalate	117-84-0	1.8e-5	<,J	Р	1.7e-5	<,J	Р	1.8e-5	<,J	P	1.6e-5	<,J	P	1.8e-5 <,J	1.9e-5	1.8e-5	Р	1.8e-5	1.7e-5
Dibenz(a,h)anthracene	53-70-3	1.9e-5	<	N	1.8e-5	<	N	1.9e-5	<,J	P	1.7e-5	<	N	1.9e-5 <,J	2.0e-5	1.9e-5	Р	1.9e-5	1.8e-5
Dibenzofuran	132-64-9	1.2e-6	<	N	1.2e-6	<	N	1.9e-6	<,J	Р	1.1e-6	<	N	1.9e-6 <,J	2.1e-6	1.9e-6	Р	1.6e-6	1.2e-6
1,2-Dichlorobenzene	95-50-1	1.2e-6	<	N	1.2e-6	<	N	1.8e-6	<,J	Р	1.1e-6	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	P	1.5e-6	1.2e-6
1,3-Dichlorobenzene	541-73-1	1.3e-6	<	N	1.3e-6	<	N	1.9e-6	<,J	Р	1.2e-6	<	N	1.9e-6 <,J	2.1e-6	1.9e-6	P	1.6e-6	1.3e-6
1,4-Dichlorobenzene	106-46-7	1.8e-6	<,J	Р	2.1e-6	J	Р	2.3e-6	J	Р	2.1e-6	<,J	P	2.3e-6 J	2.5e-6	2.3e-6	Р	2.0e-6	2.1e-6
3,3'-Dichlorobenzidine	91-94-1	1.2e-5	<	N I	1.2e-5	<	N	1.5e-5	<	N	1.1e-5	<	N	1.5e-5 <	1.6e-5	1.5e-5	N	1.3e-5	1.2e-5
2,4-Dichlorophenol	120-83-2	1.4e-6	<	N I	1.4e-6	<	N	2.0e-6	<	N	1.3e-6	<	N	2.0e-6 <	2.2e-6	2.0e-6	N N	1.7e-6	1.4e-6
Diethylphthalate	84-66-2	1.8e-6	<,J	P	1.6e-6	<	N	2.5e-6	<,J	P	1.5e-6	<	N	2.5e-6 <,J	2.7e-6	2.5e-6	P	2.1e-6	1.6e-6
Dimethyl phthalate	131-11-3	1.1e-6	<	N	1.0e-6	<	N	1.8e-6	<,J	Р	9.7e-7	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	Р	1.4e-6	1.0e-6
2,4-Dimethylphenol	105-67-9	6.5e-6	<	N	6.4e-6	<	N	7.1e-6	<	N	6.0e-6	<	N	7.1e-6 <	7.4e-6	7.1e-6	N 	6.8e-6	6.2e-6
4,6-Dinitro-2-methylphenol	534-52-1	1.4e-5	<	N	1.4e-5	<	N	1.6e-5	<	N	1.3e-5	<	N	1.6e-5 <	1.7e-5	1.6e-5	N	1.5e-5	1.4e-5
2,4-Dinitrophenol	51-28-5	3.0e-5	<	N I	2.9e-5	<	N N	3.2e-5	<	N	2.7e-5	<	N	3.2e-5 <	3.3e-5	3.2e-5	N P	3.1e-5	2.8e-5
2,4-Dinitrotoluene 2,6-Dinitrotoluene	121-14-2 606-20-2	1.4e-6 1.3e-6	<	N N	1.4e-6 1.3e-6	<	N N	2.2e-6	<,J	P	1.3e-6	<u> </u>	N N	2.2e-6 <,J	2.4e-6	2.2e-6	P	1.8e-6	1.4e-6
1,2-Diphenylhydrazine	122-66-7	1.3e-6	<	N	1.1e-6	<	N N	1.8e-6	<,J	P	1.2e-6	<		1.8e-6 <,J	2.0e-6	1.8e-6	P	1.6e-6 1.4e-6	1.3e-6
Fluoranthene	206-44-0	1.1e-6	<	N	1.1e-6	<	N	1.8e-6 1.9e-6	<,J <,J	P	9.9e-7 1.0e-6	< <	N N	1.8e-6 <,J 1.9e-6 <,J	2.0e-6 2.2e-6	1.8e-6 1.9e-6		1.4e-6 1.5e-6	1.0e-6 1.0e-6
Fluorene	86-73-7	1.1e-6	<	l N	1.1e-6	<	N	1.8e-6	۰,۰ ۲,۶	P	1.0e-6	~	N	1.8e-6 <,J	2.0e-6	1.8e-6		1.5e-6	1.0e-6
Hexachlorocyclopentadiene	77-47-4	1.8e-5	~	N	1.7e-5	~	N	2.0e-5	<u>',J</u>	P	1.6e-5		N	2.0e-5 <	2.2e-5	2.0e-5	P	1.9e-5	1.7e-5
Hexachlorobenzene	118-74-1	1.2e-6	<	N	1.2e-6	<	N I	1.8e-6	<,J	P	1.1e-6	~	N :	1.8e-6 <,J	2.0e-6	1.8e-6		1.5e-6	1.2e-6
Hexachlorobutadiene	87-68-3	1.6e-6	<	N N	1.6e-6	<	N	2.3e-6	۰,۵ <,J	P	1.5e-6	<	N	2.3e-6 <,J	2.5e-6	2.3e-6	l ' <sub>P</sub> l	2.0e-6	1.6e-6
Hexachloroethane	67-72-1	1.8e-6	<	N.	1.7e-6	<	N I	2.3e-6	<,J	P	1.6e-6	<	N	2.3e-6 <,J	2.4e-6	2.3e-6	l P	2.0e-6	1.7e-6
Indeno(1,2,3-cd)pyrene	193-39-5	1.6e-5	<	N	1.6e-5	<	N	1.7e-5	<,J	P	1.5e-5	<	N	1.7e-5 <.J	1.8e-5	1.7e-5	P	1.7e-5	1.6e-5
Isophorone	78-59-1	1.1e-6	<	N	1.1e-6	<	N	1.8e-6	<,J	P	1.0e-6	<	N	1.8e-6 <,J	2.0e-6	1.8e-6	l p l	1.5e-6	1.0e-6
2-Methylnaphthalene	91-57-6	1.1e-6	<	N	1.1e-6	<	N	1.7e-6	<,J	P	1.0e-6	<	N	1.7e-6 <,J	1.9e-6	1.7e-6	Р	1.4e-6	1.0e-6
2-Methylphenol	95-48-7	5.1e-6	<	N	5.1e-6	<	N	5.8e-6	<	N	4.8e-6	<	N	5.8e-6 <	6.1e-6	5.8e-6	N	5.5e-6	4.9e-6
3-Methylphenol & 4-Methylphenol		3.7e-6	<	N	3.7e-6	<	N	4.4e-6	<	N	3.4e-6	<	N	4.4e-6 <	4.7e-6	4.4e-6	N	4.1e-6	3.6e-6
N-Nitroso-di-n-propylamine	621-64-7	1.2e-6	<	N	1.2e-6	<	N	1.7e-6	<	N	1.1e-6	<	N	1.7e-6 <	1.8e-6	1.7e-6	N	1.5e-6	1.2e-6
N-Nitrosodimethylamine	62-75-9	1.2e-6	<	N	1.2e-6	<	N	1.7e-6	<,J	Р	1.1e-6	<	N	1.7e-6 <,J	1.8e-6	1.7e-6	P	1.5e-6	1.2e-6
N-Nitrosodiphenylamine	86-30-6	1.6e-6	<	N	1.6e-6	<	N	2.4e-6	<,J	Р	1.5e-6	<	N	2.4e-6 <,J	2.6e-6	2.4e-6	Р	2.0e-6	1.6e-6
Naphthalene	91-20-3	1.2e-6	<	N	1.2e-6	<	N	1.9e-6	<,J	P	1.1e-6	<	N	1.9e-6 <,J	2.1e-6	1.9e-6	P	1.6e-6	1.2e-6
2-Nitroaniline	88-74-4	1.2e-6	<	N	1.2e-6	<	N	1.9e-6	<	N	1.1e-6	<	N I	1.9e-6 <	2.1e-6	1.9e-6	N	1.6e-6	1.2e-6
3-Nitroaniline	99-09-2	4.5e-6	<	N	4.4e-6	<	N	5.7e-6	<	N	4.2e-6	<	N.	5.7e-6 <	6.1e-6	5.7e-6	N	5.1e-6	4.3e-6
4-Nitroaniline	100-01-6	3.9e-6	<	N	3.9e-6	<	N	5.0e-6	<	N	3.7e-6	<	N	5.0e-6 <	5.3e-6	5.0e-6	N D	4.5e-6	3.8e-6
Nitrobenzene	98-95-1	1.8e-6	<,J	P P	1.6e-6	<,J	P	2.3e-6	<,J	P P	1.2e-6	<,J	P	2.3e-6 <,J	2.6e-6	2.3e-6	P	2.0e-6	1.4e-6
2-Nitrophenol 4-Nitrophenol	88-75-5 100-02-7	1.0e-5 8.3e-6	<	P	4.4e-6 5.9e-6	<,J <	N	4.2e-6	<,J	N	5.2e-6	<,J	P P	1.0e-5 <	1.1e-5	1.0e-5	P	7.1e-6	4.8e-6
4-Nitropnenoi 2,2'-Oxybis(1-chloropropane)	100-02-7	8.3e-6 1.5e-6	<,J <	N	5.9e-6 1.5e-6	·	N	7.2e-6 2.5e-6	<	Ni P	5.6e-6 1.4e-6	<,J <	N	8.3e-6 <,J	9.2e-6	8.3e-6 2.5e-6		7.7e-6 2.0e-6	5.8e-6 1.5e-6
Pentachlorobenzene	608-93-5	1.5e-6 1.1e-6	~	N	1.1e-6	~	N	2.5e-6 1.7e-6	<,J <	N N	1.4e-6 1.0e-6	<	N N	2.5e-6 <,J 1.7e-6 <	2.8e-6 1.9e-6	2.5e-6 1.7e-6	l P	2.0e-6 1.4e-6	1.5e-6
Pentachloronitrobenzene	82-68-8	1.1e-6 1.2e-6	<	N	1.2e-6	~	N	1.7e-6 1.8e-6	<	Ň	1.1e-6	<	N N	1.7e-6 < 1.8e-6 <	2.0e-6	1.7e-6 1.8e-6	N	1.4e-6 1.5e-6	1.0e-6 1.2e-6
Pentachlorophenol	87-86-5	3.3e-5	<	N N	3.2e-5	~	N	3.5e-5	~	N	3.8e-5	<	N	3.8e-5 <	3.9e-5	3.8e-5	I N	3.4e-5	3.5e-5
Phenanthrene	85-01-8	1.1e-6	~	N	1.1e-6	~	N	1.9e-6	<,J	P	3.6 <del>e-</del> 5 9.9e-7	<	N	1.9e-6 <,J	2.2e-6	1.9e-6	P	1.5e-6	1.0e-6
Phenol	108-95-2	8.4e-6	~	P	4.0e-6	<,J	P	3.8e-6	<,J	P	4.0e-6	\ <,J	P	8.4e-6 <	9.6e-6	8.4e-6		6.1e-6	4.0e-6
Pyrene	129-00-0	1.1e-6	<	N I	1.1e-6	<	N	2.0e-6	<,J	P	1.0e-6	·,,	N	2.0e-6 <,J	2.3e-6	2.0e-6	' <sub>P</sub>	1.6e-6	1.0e-6
Pyridine	110-86-1	1.9e-6	<	N	1.8e-6	<	N	3.2e-6	<	N	1.7e-6	<	N	3.2e-6 <	3.5e-6	3.2e-6	N	2.5e-6	1.8e-6
1,2,4,5-Tetrachlorobenzene	95-94-3	1.2e-6	<	N	1.2e-6	<	N	1.8e-6	<	Ň	1.1e-6	<	Ň	1.8e-6 <	2.0e-6	1.8e-6	N	1.5e-6	1.2e-6
1,2,4-Trichlorobenzene	120-82-1	1.3e-6	<	N	1.3e-6	<	N	1.9e-6	<,J	P	1.2e-6	<	N	1.9e-6 <,J	2.1e-6	1.9e-6	P	1.6e-6	1.3e-6
2,4,5-Trichlorophenol	95-95-4	2.7e-6	<	N	2.7e-6	<	N	3.3e-6	<	N	2.5e-6	<	N	3.3e-6 <	3.5e-6	3.3e-6	N	3.0e-6	2.6e-6
2,4,6-Trichlorophenol	88-06-2	1.8e-6	<	N	1.7e-6	<	N	2.4e-6	<	N	1.6e-6	<	N	2.4e-6 <	2.6e-6	2.4e-6	N	2.1e-6	1.7e-6

Table B-17. SVOC emission rates - grames per second comparisons.

TICs	CAS Registry Number	STRT-1 g/s	Flag	Project Specific Flag	END-1 g/s	Flag	Project Specific Flag	STRT-2 g/s	Flag	Project Specific Flag	END-2 g/s	Flag	Project Specific Flag	Max value g/s	Avg+2σ g/s	Results g/s	Project Specific Flag	STRT Run Avgs g/s	END Rur Avgs g/s
1,2-Benzenedicarboxylic acid,	1330-96-7							9.6e-7	N,J,Q	Р				9.6e-7 N,J,Q		9.6e-7	Р	9.6e-7	
2,4-Hexadiene	592-46-1	l			1.7e-5	N,J,M	Р				2.8e-6	N,J,M	P	1.7e-5 N,J,M	3.1e-5	1.7e-5	P		1.0e-5
2,5-Diethylphenol	876-20-0										1.0e-5	N,J,M	Р	1.0e-5 N,J,M		1.0e-5	P		1.0e-5
2-Cyclohexene-1-one, 3-methyl-	1193-18-6										3.2e-7	N,J,M	Р	3.2e-7 N,J,M	1	3.2e-7	P	1	3.2e-7
2-Hexanone	591-78-6	2.5e-5	N,J,M	P	1.3e-5	N,J,M	Р							2.5e-5 N,J,M	3.6e-5	2.5e-5	Р	2.5e-5	1.3e-5
3-Hexanone	589-38-8	2.1e-5	N,J,M	Р	1.0e-5	N,J,M	P	1.1e-5	N,J,M	Р	7.1e-6	N,J,M	P	2.1e-5 N,J,M	2.4e-5	2.1e-5	P	1.6e-5	8.7e-6
Benzaldehyde	100-52-7	1.2e-4	N,J,M	Р	8.0e-5	N,J,M	Р	7.6e-5	N,J,M	Р	7.4e-5	N,J,M	P	1.2e-4 N,J,M	1.3e-4	1.2e-4	Р	9.8e-5	7.7e-5
Benzaldehyde, 4-ethyl-	4748-78-1				6.2e-6	N,J,M	Р							6.2e-6 N,J,M		6.2e-6	P	1 1	6.2e-6
Benzaldehyde, ethyl-	53951-50-1										6.2e-6	N,J,M	Р	6.2e-6 N,J,M		6.2e-6	Р		6.2e-6
Benzo(e)pyrene	192-97-2					Į		1.6e-6	N,J,Q	Р			1	1.6e-6 N,J,Q		1.6e-6	P	1.6e-6	
Cyclododecane	294-62-2	3.4e-6	N,J,M	Р	3.2e-6	N,J,M	Р	6.4e-7	N,J,M	Р	1.3e-6	N,J,M	Р	3.4e-6 N,J,M	4.9e-6	3.4e-6	P	2.0e-6	2.3e-6
Dodecane	112-40-3	7.3e-6	N,J,M	Р	5.2e-6	N,J,M	Р	6.3e-6	N,J,M	Р	3.8e-6	N,J,M	P	7.3e-6 N,J,M	8.7e-6	7.3e-6	l P l	6.8e-6	4.5e-6
Eicosane	112-95-8	6.7e-7	N,J,M	Р							1.9e-6	N,J,M	Р	1.9e-6 N,J,M	3.1e-6	1.9e-6	Р	6.7e-7	1.9e-6
Formic acid, phenylmethyl ester	104-57-4		, . ,	-	7.8e-6	N.J.M	Р				9.6e-6	N,J,M	P	9.6e-6 N,J,M	1.1e-5	9.6e-6	l P		8.7e-6
Furan, 2,5-dimethyl-	625-86-5	1.5e-6	N.J.M	Р	8.4e-7	N,J,M	P	1.1e-6	N,J,M	Р				1.5e-6 N,J,M	1.9e-6	1.5e-6	l p l	1.3e-6	8.4e-7
Heneicosane	629-94-7	2.2e-7	N,J,M	P		,.,.			, . ,		9.2e-7	N,J,M	Р	9.2e-7 N,J,M	1.6e-6	9.2e-7	I P I	2.2e-7	9.2e-7
Heptacosane	593-49-7										8.4e-6	N,J,M	Р	8.4e-6 N,J,M		8.4e-6	Р		8.4e-6
Heptadecane	629-78-7	9.7e-7	N,J,M	Р	4.3e-7	N,J,M	Р							9.7e-7 N.J.M	1.5e-6	9.7e-7	P	9.7e-7	4.3e-7
Heptane, 2,3-dimethyl-	3074-71-3		1	·	1.1e-6	N,J,M	P						•	1.1e-6 N,J,M		1.1e-6	l p l		1.1e-6
Heptane, 2,5-dimethyl-	2216-30-0		ł			,_,		8.1e-7	N,J,M	Р				8.1e-7 N,J,M		8.1e-7	P	4.0e-7	1
Hexacosane	630-01-3								,	-	6.5e-6	N,J,M	Р	6.5e-6 N,J,M		6.5e-6	P		6.5e-6
Hexadecanoic acid	57-10-3	5.9e-7	N,J,M	Р										5.9e-7 N,J,M		5.9e-7	l p l	5.9e-7	ŀ
Hexanedioic acid, bis(2-ethylh)	103-23-1		, . ,	·				1.1e-6	N.J.Q	P				1.1e-6 N.J.Q	Ī	1.1e-6	l P I	1.1e-6	İ
Hexatriacontane	630-06-8								,.,	-	5.8e-7	N,J,M	P	5.8e-7 N,J,M		5.8e-7	Р		5.8e-7
Naphthalene, 1-methyl-	90-12-0							8.9e-7	N,J,Q	Р		,-,		8.9e-7 N.J.Q		8.9e-7	Р	8.9e-7	
Octadecanoic acid	57-11-4	3.9e-7	N.J.M	Р	2.2e-7	N,J,M	P		, . ,					3.9e-7 N,J,M	5.6e-7	3.9e-7	Р	3.9e-7	2.2e-7
Octane, 3-methyl-	2216-33-3	8.6e-7	N,J,M	P		.,.,.	-							8.6e-7 N,J,M		8.6e-7	Р	8.6e-7	
Pentacosane	629-99-2		.,.,	·		l					3.8e-6	N,J,M	Р	3.8e-6 N,J,M		3.8e-6	P		3.8e-6
Pentadecane	629-62-9	1.6e-6	N,J,M	Р							4.6e-7	N.J.M	P	1.6e-6 N,J,M	2.7e-6	1.6e-6	Р	1.6e-6	4.6e-7
Phosphine oxide, triphenvi-	791-28-6	9.5e-7	N,J,M	P								,.,.,		9.5e-7 N,J,M		9.5e-7	P	9.5e-7	
Phosphoric acid tributyl ester	126-73-8	2.5e-6	N,J,M	P P	1.7e-6	N,J,M	Р				2.4e-6	N,J,M	Р	2.5e-6 N,J,M	3.1e-6	2.5e-6	P	2.5e-6	2.1e-6
Tetracosane	646-31-1	-::50 0	,,,,,,,	·	0 0	,,,,,,,			l		2.6e-6	N,J,M	P	2.6e-6 N,J,M	1	2.6e-6	P		2.6e-6
Tetradecane	629-59-4	9.4e-7	N.J.M	Р	1.0e-5	N,J,M	Р	6.4e-6	N,J,M	Р	1.0e-5	N.J.M	P	1.0e-5 N,J,M	1.6e-5	1.0e-5	P	3.7e-6	1.0e-5
Tetratetracontane	7098-22-8	7.2e-7	N.J.M	P		. 1,5,141	•	50 0	,0,111	·		1 1,0,111	i i	7.2e-7 N,J,M	1.00 0	7.2e-7	' <sub>P</sub>	7.2e-7	1.000
Tridecane	629-50-5	2.2e-6	N.J.M	P P	2.3e-6	N,J,M	Р	2.0e-6	N.J.M	Р	2.1e-6	N,J,M	Р	2.3e-6 N,J,M	2.3e-6	2.3e-6	P	2.1e-6	2.2e-6
Total Analytes	320 00 0		. 1,0,		2.000	. 1,0,121		1.000	. 1,0,.11			. 1,0,111	L		_,000	1.5e-3		<u> </u>	

Total Analytes
Total Detected Analytes

1.5e-3 1.2e-3 Total Volatile Emissions rate in lbs/hr = 0.011826754 Table B-18. 0031 emission rates - grams per second comparisons.

Table B-18. 0031 emission ra		per secon	d comp																			
Analysis	CAS	STRT-1		Project	END-1	⊐	Project	STRT-2	<b>T</b>	Project	END-2	<u> </u>	Project	Max value	⊐	Avg+2σ	n	Results	≖	Project	STRT Run	END Run
Analyte	Registry	g/s	Flag	Specific	g/s	Flag	Specific	g/s	Flag	Specific	g/s	Flag	Specific	g/s	Flag	g/s	Flag	g/s	Flag	Specific	Avgs g/s	Avgs
A 4	Number	00-5		Flag			Flag			Flag			Flag	20-5				20-5	D	Flag	2505	g/s
Acetone	67-64-1	2.9e-5	В	A	3.0e-5	В	A I	2.1e-5	J,B	A	1.6e-5	<,J,B	A	3.0e-5	В	3.8e-5		3.0e-5	В	A	2.5e-5	2.3e-5
Acrylonitrile	107-13-1	1.1e-5	<	N	1.2e-5	<	N	1.2e-5	<	N	1.2e-5	<	N	1.2e-5	<	1.2e-5		1.2e-5	<	N P	1.1e-5	1.2e-5
Benzene	71-43-2	1.6e-6	<	P	1.0e-6	<	P	9.1e-7	۷,>	, F	8.4e-7	<	P	1.6e-6	<	1.8e-6		1.6e-6	<	'	1.3e-6	9.3e-7
Bromobenzene	108-86-1	4.9e-7	<	N	5.0e-7	<	N	5.0e-7	<	N	5.0e-7	<	N	5.0e-7	<	5.1e-7		5.0e-7	<	N	4.9e-7	5.0e-7
Bromochloromethane	74-97-5	6.1e-7	<	N	6.3e-7	<	N	6.2e-7	<	N	6.3e-7	<	N	6.3e-7	<	6.4e-7		6.3e-7	<	N	6.2e-7	6.3e-7
Bromodichloromethane	75-27-4	4.9e-7	<	N	5.0e-7	<	N I	5.0e-7	<	N I	5.0e-7	<	N	5.0e-7	<	5.1e-7		5.0e-7	<	N	4.9e-7	5.0e-7
Bromoform	75-25-2	7.3e-7	< .	N	7.6e-7	< .	N	7.4e-7	< .	N I	7.6e-7	<	N	7.6e-7	< .	7.7e-7		7.6e-7	<	N	7.4e-7	7.6e-7
Bromomethane	74-83-9	6.9e-7	< J	P	9.7e-7	<,J	Р	6.2e-7	<,J	Р	8.4e-7	J		9.7e-7	<,J	1.1e-6		9.7e-7	<,J	P	6.6e-7	9.0e-7
2-Butanone	78-93-3	3.6e-6	<,J	P	3.7e-6	<,J	P	3.7e-6	ر,>	P	3.7e-6	< .	N	3.7e-6	<,J	3.8e-6		3.7e-6	<,J	: ' I	3.7e-6	3.7e-6
n-Butylbenzene	104-51-8	6.1e-7	<	N	6.3e-7	<	N	6.2e-7	<	N	6.3e-7	<	N	6.3e-7	<	6.4e-7		6.3e-7	<	N	6.2e-7	6.3e-7
sec-Butylbenzene	135-98-8	3.5e-7	<	N	3.6e-7	<	N I	3.5e-7	<	N	3.6e-7	<	N	3.6e-7	<	3.6e-7		3.6e-7	<	N	3.5e-7	3.6e-7
tert-Butylbenzene	98-06-6	5.7e-7	<	N	5.9e-7	<	N	5.8e-7	<	N	5.9e-7	<	N	5.9e-7	<	6.0e-7		5.9e-7	<	N	5.7e-7	5.9e-7
Carbon disulfide	75-15-0	4.0e-6	<	P	5.0e-6	<	P	5.4e-6	<	P	2.7e-6	<	P	5.4e-6	<	6.7e-6	ı	5.4e-6	<		4.7e-6	3.9e-6
Carbon tetrachloride	56-23-5	6.5e-7	<	N	6.7e-7	<	N	6.6e-7	<	N	6.7e-7	<,J		6.7e-7	<	6.8e-7		6.7e-7	<		6.6e-7	6.7e-7
Chlorobenzene	108-90-7	3.9e-7	<	N	4.0e-7	<,J	P	3.9e-7	<	N	4.0e-7	<,J	P	4.0e-7	<,J	4.1e-7		4.0e-7	<,J		3.9e-7	4.0e-7
Chlorodibromomethane	124-48-1	6.1e-7	<	N	6.3e-7	<	N	6.2e-7	< .	N	6.3e-7	<	N	6.3e-7	<	6.4e-7		6.3e-7	<	N	6.2e-7	6.3e-7
Chloroethane	75-00-3	6.5e-7	<,J	P	8.4e-7	ر,>	P	7.0e-7	<,J	Р	6.7e-7	<,J	Р	8.4e-7	<,J	8.9e-7		8.4e-7	<,J	P	6.8e-7	7.6e-7
Chloroform	67-66-3	9.8e-7	<,J	P	1.7e-6	<	P	1.4e-6	<	Р	1.6e-6	<	Р	1.7e-6	<	2.1e-6		1.7e-6	<	P	1.2e-6	1.7e-6
Chloromethane	74-87-3	4.9e-6	J	P	1.2e-5	<	P	5.0e-6	ا,>	P	1.1e-5	<,J	Р	1.2e-5	<	1.6e-5	ŀ	1.2e-5	<		4.9e-6	1.1e-5
2-Chlorotoluene	95-49-8	2.4e-7	<	N N	2.5e-7	<	N	2.4e-7	<	N	2.5e-7	<	N	2.5e-7	<	2.5e-7		2.5e-7	<	N N	2.4e-7	2.5e-7
4-Chlorotoluene	106-43-4	2.4e-7	<	N	2.5e-7	<	N I	2.4e-7	<	N I	2.5e-7	<	N	2.5e-7	<	2.5e-7		2.5e-7	<	N	2.4e-7	2.5e-7
1,2-Dibromo-3-chloropropane	96-12-8	1.1e-6	<	N N	1.2e-6	<	N	1.2e-6	<	N N	1.1e-6	<	N	1.2e-6	<	1.2e-6		1.2e-6	<	N	1.1e-6	1.2e-6
1,2-Dibromoethane	106-93-4	8.2e-7	<	N I	8.4e-7	<	N	8.3e-7	<	N	8.4e-7	<	N	8.4e-7	<	8.6e-7		8.4e-7	<	N	8.2e-7	8.4e-7
Dibromomethane	74-95-3	6.9e-7	<	N N	7.2e-7	<	N	7.0e-7	<	N	7.1e-7	<	N	7.2e-7	<	7.3e-7		7.2e-7	<	N	7.0e-7	7.2e-7
1,2-Dichlorobenzene	95-50-1	7.3e-7	<	N	7.6e-7	<	N	7.4e-7	<	N	7.6e-7	<	N	7.6e-7	<	7.7e-7	l	7.6e-7	<	N	7.4e-7	7.6e-7
1,3-Dichlorobenzene	541-73-1	4.1e-7	<	N	4.2e-7	<	N	4.1e-7	<	N	4.2e-7	<	N	4.2e-7	<	4.3e-7	I	4.2e-7	<	N	4.1e-7	4.2e-7
1,4-Dichlorobenzene	106-46-7	5.7e-7	<	N	5.9e-7	< .	N	5.8e-7	<	N	5.9e-7	<.	N <sub>D</sub>	5.9e-7	<	6.0e-7	I	5.9e-7	<	N	5.7e-7	5.9e-7
Dichlorodifluoromethane	75-71-8	1.1e-6	<	P	6.7e-7	<,J	P	8.3e-7	<	P	6.7e-7	<,J	1	1.1e-6	<	1.2e-6		1.1e-6	<	P	9.4e-7	6.7e-7
1,1-Dichloroethane	75-34-3	6.1e-7	<	N	6.3e-7	<	N	6.2e-7	<	N	6.3e-7	<	N	6.3e-7	<	6.4e-7	l	6.3e-7	<	N P	6.2e-7	6.3e-7
1,2-Dichloroethane	107-06-2	6.5e-7	<,J	P	6.7e-7	<,J	Р	6.6e-7	< .	N	6.7e-7	<.	N	6.7e-7	<,J	6.8e-7		6.7e-7	<,J		6.6e-7	6.7e-7
1,1-Dichloroethene	75-35-4	6.5e-7	<,J	P	7.6e-7	ل,>	P	6.6e-7	<,J	P	6.7e-7	<,J	P	7.6e-7	<,J	7.8e-7		7.6e-7	<,J	' '	6.6e-7	7.2e-7
cis-1,2-Dichloroethene	156-59-2	6.1e-7	<	N	6.3e-7	<	N	6.2e-7	<	N	6.3e-7	<	N	6.3e-7	<	6.4e-7		6.3e-7	<	N	6.2e-7	6.3e-7
trans-1,2-Dichloroethene	156-60-5	6.9e-7	<	N I	6.7e-7	< .	N	6.6e-7	<	N I	7.1e-7	<	N	7.1e-7	<	7.3e-7		7.1e-7	<	N D	6.8e-7	6.9e-7
1,2-Dichloropropane	78-87-5	5.3e-7	<	N I	5.5e-7	<,J	P	5.4e-7	<	N	5.5e-7	<	N	5.5e-7	<,J	5.6e-7		5.5e-7	<,J	' '	5.3e-7	5.5e-7
1,3-Dichloropropane	142-28-9	6.9e-7	<	N I	7.2e-7	<	N	7.0e-7	<	N	7.1e-7	<	N	7.2e-7	<	7.3e-7		7.2e-7	<	N I	7.0e-7	7.2e-7
2,2-Dichloropropane	594-20-7	6.5e-7	<	N	6.7e-7	<	N	6.6e-7	<	N	6.7e-7	<	N	6.7e-7	<	6.8e-7		6.7e-7	<	N	6.6e-7	6.7e-7
1,1-Dichloropropene	563-58-6	7.3e-7	<	N	7.6e-7	<	N	7.4e-7	<	N	7.6e-7	<	N	7.6e-7	<	7.7e-7	- 1	7.6e-7	<	N I	7.4e-7	7.6e-7
cis-1,3-Dichloropropene	10061-01-5	5.3e-7	<	N	5.5e-7	<	N	5.4e-7	<	N	5.5e-7	<	N	5.5e-7	<	5.6e-7	l	5.5e-7	<	N I	5.3e-7	5.5e-7
trans-1,3-Dichloropropene	10061-02-6	6.1e-7	<	N	6.3e-7	<	N	6.2e-7	<	N	6.3e-7	<	N	6.3e-7	<	6.4e-7	ľ	6.3e-7	<	N	6.2e-7	6.3e-7
Ethylbenzene	100-41-4	3.8e-7	<	N	3.9e-7	<	N N	3.8e-7	<	N	3.9e-7	<	N	3.9e-7	<	4.0e-7		3.9e-7	<	N N	3.8e-7	3.9e-7
Hexachlorobutadiene	87-68-3	9.0e-7	<	N	9.3e-7	<	N	9.1e-7	<	N	9.7e-7	<	N	9.7e-7	<	9.9e-7		9.7e-7	<	N	9.0e-7	9.5e-7
2-Hexanone	591-78-6	2.3e-6	<	N	2.4e-6	<	N	2.3e-6	<	N	2.4e-6	<	N	2.4e-6	<	2.4e-6		2.4e-6	<	N	2.3e-6	2.4e-6
Isopropylbenzene	98-82-8	2.9e-7	<	N	2.9e-7	<	N	2.9e-7	<	N	2.9e-7	<	N	2.9e-7	<	3.0e-7		2.9e-7	<	N	2.9e-7	2.9e-7
p-Isopropyltoluene	99-87-6	4.5e-7	F B	N	4.6e-7	< <u> </u>	N	4.5e-7	< D	N A	4.6e-7	< ID	N A	4.6e-7	< E D	4.7e-7	$\rightarrow$	4.6e-7	< P	N N	4.5e-7	4.6e-7 4.5e-6
Methylene chloride	75-09-2	7.3e-5	_,_	A	7.2e-6	В	A	4.5e-6	<,B	A	1.8e-6	<,J,B	A		E,B	9.1e-5	J	7.3e-5	E,B	A	3.9e-5	4.5e-6 2.5e-6
4-Methyl-2-pentanone	108-10-1	2.4e-6	<	N	2.5e-6	<	N	2.4e-6	<	N	2.5e-6	<	N	2.5e-6	<	2.5e-6	J	2.5e-6	<	N	2.4e-6	2.5e-6 8.8e-7
Naphthalene	91-20-3	8.6e-7	<	N N	8.8e-7	< <	N	8.7e-7	<	N N	8.8e-7	<	N	8.8e-7	<	9.0e-7	J	8.8e-7	< <	N	8.6e-7	2.8e-7
n-Propylbenzene	103-65-1	2.7e-7		N N	2.8e-7	<	N N	2.7e-7	<	N N	2.8e-7	< <	N N	2.8e-7	<	2.8e-7		2.8e-7 3.3e-7	- <	N N	2.7e-7 3.2e-7	3.3e-7
Styrene	100-42-5 630-20-6	3.2e-7	<	N	3.3e-7	<	N N	3.2e-7	<		3.3e-7	- 1	N N	3.3e-7	<	3.3e-7 4.3e-7	- 1	3.3e-7 4.2e-7	<	N	3.2e-7 4.1e-7	3.3e-7 4.2e-7
1,1,1,2-Tetrachloroethane		4.0e-7	<	N	4.2e-7	<	N N	4.1e-7		N N	4.2e-7	<		4.2e-7	<			4.2e-7 9.7e-7	`	N	4.1e-7 9.0e-7	4.2e-7 9.5e-7
1,1,2,2-Tetrachloroethane	79-34-5	9.0e-7	<	N N	9.3e-7		N	9.1e-7	< <	N N	9.7e-7	<u> </u>	N	9.7e-7		9.9e-7				N	9.0e-7 6.2e-7	9.5e-7 6.3e-7
Tetrachloroethene	127-18-4	6.1e-7	<	N	6.3e-7	<	N	6.2e-7		N P	6.3e-7	<	N P	6.3e-7	<	6.4e-7		6.3e-7	<	P		9.7e-7
Toluene	108-88-3	7.8e-7	<,J	P	1.2e-6	<,J	Р	1.2e-6	<,J		7.6e-7	<,J			<,J	1.5e-6		1.2e-6	<,j		1.0e-6	
1,2,3-Trichlorobenzene	87-61-6	8.6e-7	<	N	8.8e-7	<	N	8.7e-7	< <	N	8.8e-7	<u> </u>	N N	8.8e-7	<	9.0e-7 9.9e-7		8.8e-7	< <	N N	8.6e-7 9.0e-7	8.8e-7 9.5e-7
1,2,4-Trichlorobenzene	120-82-1	9.0e-7	<	N	9.3e-7	<	N	9.1e-7	<	N N	9.7e-7	<u> </u>	N	9.7e-7	< <	B .	J	9.7e-7	·	N N	9.0e-7 7.8e-7	9.5e-7 8.0e-7
1,1,1-Trichloroethane	71-55-6	7.8e-7		N N	8.0e-7	<	N N	7.8e-7		N N	8.0e-7	<	N N	8.0e-7		8.1e-7		8.0e-7		N N	7.8e-7 7.0e-7	
1,1,2-Trichloroethane	79-00-5	6.9e-7	<	N N	7.2e-7	<	N	7.0e-7	<	N	7.1e-7	<	N	7.2e-7	<	7.3e-7		7.2e-7	<	N N		7.2e-7
Trichloroethene	79-01-6	6.5e-7	<	N	6.7e-7	۲ .	N	6.6e-7	<	N	6.7e-7	<u> </u>	N	6.7e-7	<	6.8e-7		6.7e-7	<	N	6.6e-7	6.7e-7
Trichlorofluoromethane	75-69-4	6.5e-7	<,J	P	6.7e-7	<,J	P	6.6e-7	<,J	Р	6.7e-7	<,J	Р		<,J	6.8e-7	1	6.7e-7	<,J	P	6.6e-7	6.7e-7
1,2,3-Trichloropropane	96-18-4	9.8e-7	<	N	1.0e-6	<	N	9.9e-7	<	N N	1.0e-6	<	N.	1.0e-6	<	1.0e-6		1.0e-6	<	N.	9.8e-7	1.0e-6
1,2,4-Trimethylbenzene	95-63-6	4.1e-7	<	N	4.2e-7	<	N	4.1e-7	<	N	4.2e-7	<	N	4.2e-7	<	4.3e-7	J	4.2e-7	<	N N	4.1e-7	4.2e-7
1,3,5-Trimethylbenzene	108-67-8	2.4e-7	<	N	2.5e-7	<	N	2.4e-7	<	N	2.5e-7	<	N	2.5e-7	<	2.5e-7	-	2.5e-7	<	N	2.4e-7	2.5e-7
Vinyl chloride	75-01-4	5.3e-7	<,J	P	7.6e-7	<,J	P	5.4e-7	<,J	P	7.1e-7	<,J	P		<,J	8.7e-7	1	7.6e-7	<,J	P	5.3e-7	7.4e-7
m-Xylene & p-Xylene	36777-61-	1.8e-6	<	N	1.9e-6	<	N	1.8e-6	<	N	1.9e-6	<	N D	1.9e-6	<	1.9e-6		1.9e-6	<	N	1.8e-6	1.9e-6
o-Xylene	95-47-6	3.1e-7	<	N	3.3e-7	<,J	Р	3.2e-7	<	N	3.2e-7	<,J	Р	3.3e-7	<,J	3.3e-7		3.3e-7	<,J	Р	3.2e-7	3.3e-7

Table B-18. 0031 emission rates - grams per second comparisons.

TICs	CAS Registry Number	STRT-1 g/s	Flag	Project Specific Flag	END-1 g/s	Flag	Project Specific Flag	STRT-2 g/s	Flag	Project Specific Flag	END-2 g/s	Flag	Project Specific Flag	Max value g/s	Flag	Avg+2σ g/s	Flag	Results g/s	Flag	Project Specific Flag	STRT Run Avgs µg/dscm	END Run Avgs μg/dscm
1-Heptene	592-76-7	2.2e-7	N,J,M	Р										2.2e-7	N,J,M			2.2e-7	N,J,M	Р	2.2e-7	
Benzonitrile	100-47-0				8.0e-7	N,J,M	Р	3.0e-7	N,J,M	Р	2.0e-7	N,J,M	Р	8.0e-7	N,J,M	1.1e-6		8.0e-7	N,J,M	Р	3.0e-7	5.0e-7
Butane, 1-chloro-	109-69-3	2.3e-7	N,J,M	Р										2.3e-7	N,J,M			2.3e-7	N,J,M	Р	2.3e-7	1
Cyclobutane, ethenyl-	2597-49-1				5.0e-7	N,J,M	Р							5.0e-7	N,J,M			5.0e-7	N,J,M	Р		5.0e-7
Cyclohexane, hexyl-	4292-75-5	2.4e-7	N,J,M	Р										2.4e-7	N,J,M			2.4e-7	N,J,M	Р	2.4e-7	
Cyclohexane, methyl-	108-87-2	4.5e-7	N,J,M	Р	8.4e-7	N,J,M	Р	8.3e-7	N,J,M	Р	5.0e-7	N,J,M	Р	8.4e-7	N,J,M	1.1e-6		8.4e-7	N,J,M	Р	6.4e-7	6.7e-7
Cyclohexene	110-83-8	4.1e-7	N,J,M	Р	1.5e-7	N,J,M	Р	7.4e-7	N,J,M	Р	1.1e-7	N,J,M	Р	7.4e-7	N,J,M	9.4e-7		7.4e-7	N,J,M	Р	5.8e-7	1.3e-7
Cyclopentane, 1,2-dimethyl-	2452-99-5							2.2e-7	N,J,M	Р				2.2e-7	N,J,M		ľ	2.2e-7	N,J,M	Р	2.2e-7	
Cyclopentane, 1,2-dimethyl-, t	822-50-4				3.3e-7	N,J,M	Р				1.3e-7	N,J,M	Р	3.3e-7	N,J,M	5.2e-7		3.3e-7	N,J,M	Р		2.3e-7
Cyclopentane, ethyl-	1640-89-7	1.1e-7	N,J,M	Р	1.7e-7	N,J,M	Р	1.5e-7	N,J,M	Р		İ		1.7e-7	N,J,M	2.0e-7	1	1.7e-7	N,J,M	Р	1.3e-7	1.7e-7
Decane	124-18-5	4.9e-7	N,J,M	Р										4.9e-7	N,J,M			4.9e-7	N,J,M	Р	4.9e-7	1
Decane, 2,2,5-trimethyl-	62237-96-1							2.5e-7	N,J,M	Р				2.5e-7	N,J,M			2.5e-7	N,J,M	Р	2.5e-7	i
Decane, 2,9-dimethyl-	1002-17-1	2.6e-7	N,J,M	Р										2.6e-7	N,J,M			2.6e-7	N,J,M	Р	2.6e-7	
Dodecane	112-40-3	1.3e-4	N,J,M	Р	1.1e-4	N,J,M	Р	9.1e-5	N,J,M	Р	7.1e-5	N,J,M	Р	1.3e-4	N,J,M	1.5e-4		1.3e-4	N,J,M	Р	1.1e-4	9.0e-5
Dodecane, 6-methyl-	6044-71-9							2.1e-7	N,J,M	Р				2.1e-7	N,J,M		ı	2.1e-7	N,J,M	Р	2.1e-7	
Hexadecane	544-76-3				8.8e-7	N,J,M	Р					Ī		8.8e-7	N,J,M		ł	8.8e-7	N,J,M	Р	1 1	8.8e-7
Hexane, 2,4-dimethyl-	589-43-5	4.5e-7	N,J,M	Р	7.6e-7	N,J,M	Р	8.3e-7	N,J,M	Р	4.0e-7	N,J,M	Р	8.3e-7	N,J,M	1.0e-6		8.3e-7	N,J,M	Р	6.4e-7	5.8e-7
Hexane, 2-methyl-	591-76-4	6.9e-7	N,J,M	P	1.3e-6	N,J,M	Р	1.2e-6	N,J,M	Р	1.0e-6	N,J,M	Р	1.3e-6	N,J,M	1.6e-6		1.3e-6	N,J,M	Р	9.7e-7	1.2e-6
Hexane, 3-methyl-	589-34-4	1.6e-6	N,J,M	Р	3.2e-6	N,J,M	Р	1.1e-6	N,J,M	Р	1.4e-6	N,J,M	Р	3.2e-6	N,J,M	3.7e-6		3.2e-6	N,J,M	Р	1.3e-6	2.3e-6
Methane, trichloronitro-	76-06-2				1.5e-6	N,J,M	Р							1.5e-6	N,J,M			1.5e-6	N,J,M	Р	1 1	1.5e-6
Octane	111-65-9	1.1e-7	N,J,M	Р										1.1e-7	N,J,M			1.1e-7	N,J,M	Р	1.1e-7	
Pentane, 2,3-dimethyl-	565-59-3	7.3e-7	N,J,M	Р	1.3e-7	N,J,M	Р	4.5e-7	N,J,M	Р	5.9e-7	N,J,M	Р	7.3e-7	N,J,M	9.9e-7		7.3e-7	N,J,M	Р	5.9e-7	3.6e-7
Pentane, 3,3-dimethyl-	562-49-2				2.5e-7	N,J,M	Р							2.5e-7	N,J,M			2.5e-7	N,J,M	Р	1 1	2.5e-7
Pentane, 3-ethyl-	617-78-7					·		1.4e-7	N,J,M	Р	1.3e-7	М,Ј,М	Р	1.4e-7	N,J,M	1.5e-7	- 1	1.4e-7	N,J,M	Р	1.4e-7	1.3e-7
Tetradecane	629-59-4	3.8e-6	N,J,M	Р	4.2e-6	N,J,M	Р	4.0e-6	N,J,M	Р	5.9e-6	N.J.M	Р	5.9e-6	N.J.M	6.4e-6		5.9e-6	N,J,M	Р	3.9e-6	5.0e-6
Tridecane	629-50-5	1.1e-5	N,J,M	Р	1.3e-5	N,J,M	P	3.4e-7	N,J,M	Р	1.3e-5	N,J,M	P		N,J,M	2.2e-5		1.3e-5	N,J,M	Р	5.9e-6	1.3e-5
Undecane	1120-21-4	4.1e-6	N,J,M	Р	2.1e-6	N,J,M	P	9.9e-7	N,J,M	Р	5.0e-7	N,J,M	Р	4.1e-6	N,J,M	5.1e-6	1	4.1e-6	N,J,M	Р	2.5e-6	1.3e-6
Undecane, 2,6-dimethyl-	17301-23-4	4.5e-7	N,J,M	Р				1.9e-7	N,J,M	Р	3.5e-7	N,J,M	Р	4.5e-7	N,J,M	5.9e-7	ĺ	4.5e-7	N,J,M	P	3.2e-7	3.5e-7
Undecane, 5-methyl-	1632-70-8	2.6e-6	N,J,M	Р	7.6e-7	N,J,M	Р	8.7e-7	N,J,M	Р	7.6e-7	N,J,M	Р		N,J,M	3.1e-6		2.6e-6	N,J,M	Р	1.7e-6	7.6e-7
Total Analytes Total Detected Analytes									· · · · · · · · · · · · · · · · · · ·			1						3.5e-4 3.0e-4				

Total Semivolatiles in lbs/hr

0.002769813

Table B-19. 0050 emission rates - grams per second comparisons.

Analyte	STRT-1 g/s	Flag	END-1 g/s	Flag	STRT-2 g/s	Flag	END-2 g/s	Flag	Max value	Flag	Avg+2o <u>T</u> g/s ©	Results g/s	Flag	STRT Run Avgs g/s	END Run Avgs g/s
Chloride (as HCI)	3.3e-4	В	2.9e-4	В	3.2e-4	В	3.2e-4	В	3.3e-4	В	3.4e-4	3.3e-4	В	3.2e-4	3.1e-4
Chloride (as CI2)	3.3e-5	<	5.8e-5	В	3.0e-5	<	3.2e-5	<	5.8e-5	В	6.4e-5	5.8e-5	В	3.2 <b>e</b> -5	4.5e-5
Fluoride (as HF)	4.5e-5	<	4.0e-5	<	4.1e-5	<	4.3e-5	<	4.5e-5	<	4.7e-5	4.5e-5	<	4.3e-5	4.2e-5
Nitrate (as HNO3)	1.4e-3		1.1e-3		9.4e-4		7.9e-4		1.4e-3		1.5e-3	1.4e-3	ŀ	1.1e-3	9.2e-4
Nitrite (as HNO2)	1.8e-4	<	3.8e-4	<	1.7e-4	<	3.4e-4	<	3.8e-4	<	4.9e-4	3.8e-4	<	1.8e-4	3.6e-4
Particulate	4.0e-4	В	2.9e-4	В	1.2e-4		8.9e-5	В	4.0e-4	В	5.2e-4	4.0e-4	В	2.6e-4	1.9e-4

maximum hourly emissions

Table B-20. 0060 emission rates - grams per second comparisons.

Analyte	CAS Registry Number	STRT-1 g/s	Flag	Project Specific Flags	END-1 g/s	Flag	Project Specific Flags	STRT-2 g/s	Flag	Project Specific Flags	END-2 g/s	Flag	Project Specific Flags	Max value g/s	Flag	Avg+2σ g/s	Flag	Results g/s	Flag	Project Specific Flag	STRT Run Avgs g/s	END Run Avgs g/s
Aluminum (Al)	7429-90-5	1.6e-5		Α	2.5e-5		Α	8.3e-6		Α	8.2e-6		Α	2.5e-5		3.4e-5		2.5e-5		Α	1.2e-5	1.7e-5
Antimony (Sb)	7440-36-0	5.3e-7、	В	Α	4.8e-7	В	A	4.9e-7	В	A	4.6e-7	В	A	5.3e-7	В	5.5e-7		5.3e-7		A	5.1e-7	4.7e-7
Arsenic (As)	7440-38-2	1.8e-7	<,B	Р	1.5e-7	<	N I	1.8e-7	<	N	1.8e-7	<	N	1.8e-7	<,B	2.0e-7		1.8e-7		P	1.8e-7	1.6e-7
Barium (Ba)	7440-39-3	7.7 <b>e</b> -7	В	Α	9.4e-7	В	A	5.6e-7	В	A	5.3e-7	В	Α	9.4 <b>e-</b> 7	В	1.1e-6		9.4e-7		Α	6.6e-7	7.3e-7
Beryllium (Be)	7440-41-7	7.4e-8	<,B	Р	6.1e-8	<,B	Р	7.1e-8	<,B	Р	7.3e-8	<,B	Р	7.4e-8	ν,Β	8.3e-8		7.4e-8		Р	7.3e-8	6.7e-8
Cadmium (Cd)	7440-43-9	5.3e-8	В	A	7.7e-8	В	A	3.0e-8	<,B	P	3.0e-8	<	N	7.7e-8	В	1.0e-7		7.7e-8		Р	4.1e-8	5.3e-8
Chromium (Cr)	7440-47-3	3.4e-7		A	4.4e-7		A	2.7e-7		A	5.7e-7		A	5.7e-7		5.2e-7	1	5.2e-7		Α	3.1e-7	5.1e-7
Cobalt (Co)	7440-48-4	3.1e-7	В	Α	2.5e-7	<	N	3.0e-7	<	N	3.0e-7	<	N	3.1e-7	В	3.5e-7	1	3.1e-7		Р	3.0e-7	2.8e-7
Copper (Cu)	7440-50-8	4.9e-7	В	Α	3.4e-7	В	Α	2.5e-7	В	A	1.6e-7	<,B	P	4.9e-7	В	6.0e-7		4.9e-7		Р	3.7e-7	2.5e-7
Lead (Pb)	7439-92-1	1.5e-7	<,B	P	1.2e-7	В	Α	1.5e-7	<,B	P	1.3e-7	<,B	P	1.5e-7	<,B	1.8e-7		1.5e-7		Р	1.5e-7	1.3e-7
Manganese (Mn)	7439-96-5	2.2e-6		A	2.2e-6		Α	4.2e-6		Α	8.0e-6		Α	8.0e-6		5.1e-6		5.1e-6		Α	3.2e-6	5.1e-6
Mercury (Hg)	7439-97-6	1.1e-5		A	1.4e-5		Р	1.2e-5		Р	1.2e-5		P	1.4e-5		1.5e-5		1.4e-5		Р	1.2e-5	1.3e-5
Nickel (Ni)	7440-02-0	5.7e-7	В	Α	5.6e-7	В	Α	4.9e-7	В	Α	4.6e-7	В	Α	5.7e-7	В	6.3e-7		5.7e-7		Α	5.3e-7	5.1e-7
Selenium (Se)	7782-49-2	3.7e-7	В	A	2.0e-7	<	Р	2.9e-7	В	Α	2.6e-7	В	Α	3.7e-7	В	4.5e-7		3.7e-7		P	3.3e-7	2.3e-7
Silver (Ag)	7440-22-4	3.0e-7	<	N	2.4e-7	<	N	2.9e-7	<	N	2.9e-7	<	N	3.0e-7	<	3.3e-7		3.0e-7		N	2.9e-7	2.7e-7
Thallium (TI)	7440-28-0	2.9e-7	<	N	2.2e-7	<	N	2.7e-7	<	N	2.7e-7	<	N	2.9e-7	<	3.3e-7		2.9e-7		N	2.8e-7	2.5e-7
Vanadium (V)	7440-62-2	3.1e-7	<	N	2.5e-7	<	N	3.0e-7	<	N	3.0e-7	<	N	3.1 <b>e</b> -7	<	3.5e-7		3.1e-7			3.0e-7	2.8e-7
Zinc (Zn)	7440-66-6	7.0e-6		Α	1.1e-5		Α	3.3e-6		Α	1.9e-6		Α	1.1e-5		1.5e-5		1.1e-5		A	5.1e-6	6.6e-6
Total Metals														-				6.1e-5				

6.0e-5

Total Detected Metals

Table B-21. 0060 blank corrected emission rates - grams per second comparisons.

Analyte	CAS Registry Number	STRT-1 g/s	Flag	Project Specific Flags	END-1 g/s	Flag	Project Specific Flags	STRT-2 g/s	Flag	Project Specific Flags	END-2 g/s	Flag	Project Specific Flags		Flag	Avg+2σ g/s	Flag	Results g/s	Flag	Project Specific Flag	STRT Run Avgs g/s	END Run Avgs g/s
Aluminum (Al)	7429-90-5	1.5e-5		Α	2.4e-5		Α	7.5e-6		Α	7.5e-6		Α	2.4e-5		3.2e-5		2.4e-5		A	1.1e-5	1.6e-5
Antimony (Sb)	7440-36-0	1.2e-7	В	A	1.4e-7	В	Α	1.0e-7	В	Α	1.0e-7	В	Α	1.4e-7	В	1.6e-7		1.4e-7		Α	1.1e-7	1.2e-7
Arsenic (As)	7440-38-2	9.9e-8	<,B	P	8.1e-8	<	N	9.4e-8	<	N	9.4e-8	<	N	9.9e-8	<,B	1.1e-7		9.9e-8		Р	9.6e-8	8.7e-8
Barium (Ba)	7440-39-3	2.5e-7	В	Α	5.2e-7	В	Α	5.9e-8	В	Α	5.9e <b>-8</b>	В	A	5.2e-7	В	7.4e-7		5.2e-7		A	1.5e-7	2.9e-7
Beryllium (Be)	7440-41-7	5.2e-8	<,B	P	4.2e-8	<,B	Р	4.9e-8	<,B	Р	4.9e-8	<,B	Р	5.2e-8	<,B	5.8e-8		5.2e-8		Р	5.0e-8	4.6e-8
Cadmium (Cd)	7440-43-9	4.6e-8	В	A	7.1e-8	В	Α	2.3e-8	<,B	Р	2.3e-8	<,B	N	7.1e-8	В	9.5e-8		7.1e-8		Р	3.5e-8	4.7e-8
Chromium (Cr)	7440-47-3	1.3e-7		A	2.7e-7		Α :	7.4e-8		Α	7.4e-8		Α	2.7e-7		3.6e-7		2.7e-7		A	1.0e-7	1.7e-7
Cobalt (Co)	7440-48-4	3.1e-7	В	Α	2.5e-7	<	N	3.0e-7	<	N	3.0e-7	<	N	3.1e-7	В	3.5e-7		3.1e-7		Р	3.0e-7	2.7e-7
Copper (Cu)	7440-50-8	5.0e-7	В	Α	3.4e-7	В	Α	2.5e-7	В	Α	2.5e-7	В	Р	5.0e-7	В	6.1e-7		5.0e-7		Р	3.7e-7	2.9e-7
Lead (Pb)	7439-92-1	1.0e-7	<,B	P	8.2e-8	В	Α	1.1e-7	<,B	P	1.1e-7	<,B	Р	1.1e-7	<,B	1.2e-7		1.1e-7		P	1.0e-7	9.5e-8
Manganese (Mn)	7439-96-5	2.0e-6		Α	2.0e-6		Α	4.0e-6		Α	4.0e-6		Α	4.0e-6		5.0e-6		4.0e-6		Α	3.0e-6	3.0e-6
Mercury (Hg)	7439-97-6	1.1e-5		Α	1.4e-5		P	1.2e-5		Р	1.2e-5		P	1.4e-5		1.5e-5		1.4e-5		P	1.2e-5	1.3e-5
Nickel (Ni)	7440-02-0	1.8e-7	В	Α	2.4e-7	В	Α	1.1e-7	В	Α	1.1e-7	В	Α	2.4e-7	В	3.0e-7		2.4e-7		Α	1.5e-7	1.8e-7
Selenium (Se)	7782-49-2	1.1e-17	В	A	9.4e-18	<	Р	1.1e-17	В	A	1.1e-17	В	Α	1.1e-17	В	1.3e-17		1.1e-17		P	1.1e-17	1.0e-17
Silver (Ag)	7440-22-4	3.0e-7	<	N	2.4e-7	<	N	2.9e-7	<	N	2.9e-7	<	N	3.0e-7	<	3.3e-7		3.0e-7		N	2.9e-7	2.7e-7
Thallium (TI)	7440-28-0	2.9e-7	<	N	2.2e-7	<	N	2.7e-7	<	N	2.7e-7	<	N	2.9e-7	<	3.3e-7		2.9e-7		N	2.8e-7	2.5e-7
Vanadium (V)	7440-62-2	3.1e-7	<	N	2.5e-7	<	N	3.0e-7	<	N	3.0e-7	<	N	3.1e-7	<	3.5e-7		3.1e-7		N	3.0e-7	2.7e-7
Zinc (Zn)	7440-66-6	6.4e-6		Α	1.1e-5		Α	2.8e-6		Α	2.8e-6		Α	1.1e-5		1.5e-5		1.1e-5		A	4.6e-6	6.8e-6
Total Metals														*****				5.6e-5		•	<u> </u>	

5.5e-5

Total Detected Metals

Table B-22. SVOC concentration-basis.

Table B-22. SVOC concentration-				L Project I			1 Droinet			I Droinet			Uroject			· · · · · · · · · · · · · · · · · · ·			Urorect	STRIRUM	END Run
Analyte	CAS Registry	STRT-1	Flag	Project Specific	END-1	Flag	Specific	STRT-2	Flag	Project Specific	END-2	Flag	Project Specific	Max value	Flag	Avg+2σ =	Results	고	Project Specific	Avgs	Avgs
Analyte	Number	μg/dscm	99	Flag	μg/dscm	Đ	Flag	μg/dscm	g	Flag	μg/dscm	ag	Flag	μg/dscm	Ö	μg/dscm 🗳	μg/dscm	ag	Flag	μg/dscm	μg/dscm
Acenaphthene	83-32-9	3.1e0	<	N	3.2e0	<	N	5.2e0	<,J	P	3.0e0	<	N	5.2e0	<,J	5.7e0	5.2e0	$\dashv$	P	4.2e0	3.1e0
Acenaphthylene	208-96-8	3.0e0	<	N	3.1e0	<	l 'n	5.2e0 5.2e0	۰,۵ <,J	P	2.9e0	<	N	5.2e0	-,5 <,J	5.7e0 5.7e0	5.2e0 5.2e0	ı	'p	4.1e0	3.0e0
Acetophenone	98-86-2	1.6e1	<,J	'P	1.2e1	<,J	'P	1.4e1	<,J	l 'P	1.1e1	<,J	P	1.6e1	<,J	1.7e1	1.6e1	ł	P	1.5e1	1.2e1
Aniline	62-53-3	3.5e1	<	l N	3.6e1	<	N	4.8e1	<	N	3.4e1	<,0	N	4.8e1	<	5.2e1	4.8e1	- 1	N	4.2e1	3.5e1
Anthracene	120-12-7	3.0e0	<u> </u>	N	3.1e0	<	i N	5.2e0	<,J	P	2.9e0		N	5.2e0	<,J	5.7e0	5.2e0		P	4.1e0	3.0e0
Benzidine	92-87-5	2.1e2	<	l N	2.2e2	<	N	2.6e2	<	N	2.1e2	<	N N	2.6e2	<	2.7e2	2.6e2		N	2.4e2	2.1e2
Benzoic acid	65-85-0	1.6e3	E	P	8.5e2	<,E	P	7.4e2	E	P	8.3e2	Ē	P	1.6e3	Ē	1.8e3	1.6e3		P	1.2e3	8.4e2
Benzo(a)anthracene	56-55-3	3.8e0	_	N	3.9e0	<	l 'n	6.1e0	- <,J	P	3.7e0	<	N	6.1e0	<,J	6.7e0	6.1e0		P	5.0e0	3.8e0
Benzo(a)pyrene	50-32-8	4.2e1	<del>-</del>	N	4.3e1	· <	N	4.5e1	<,J	<del>'</del>	4.0e1	~~	N	4.5e1	<,J	4.7e1	4.5e1	$\dashv$	P	4.3e1	4.1e1
Benzo(b)fluoranthene	205-99-2	9.6e1	<	l N	9.8e1	<	N	9.7e1	۰,۵ ۷,۶	P	9.2e1	<	N	9.8e1	<	1.0e2	9.8e1	- 1	P	9.6e1	9.5e1
Benzo(g,h,i)perylene	191-24-2	5.4e1	<	N	5.6e1	<	N N	5.8e1	<,J	P	5.2e1	<	N	5.8e1	<,J	6.0e1	5.8e1	ı	P	5.6e1	5.4e1
Benzo(k)fluoranthene	207-08-9	1.4e2	<	N	1.4e2	<	N	1.4e2	۰,۵ <,J	P	1.3e2	<	N	1.4e2	<	1.4e2	1.4e2		P	1.4e2	1.4e2
Benzyl alcohol	100-51-6	1.8e2	<	N	1.9e2	<	N	1.8e2	<	N	1.8e2	<del>-</del>	N	1.9e2	<u> </u>	1.9e2	1.9e2		<u>i</u>	1.8e2	1.8e2
bis(2-Chloroethoxy)methane	111-91-1	3.2e0	<	N	3.3e0	<	N	4.5e0	<	N	3.1e0	<	l n	4.5e0	<	4.9e0	4.5e0		Ň	3.9e0	3.2e0
bis(2-Chloroethyl)ether	111-44-4	3.5e0	<	N	3.6e0	<	N	5.2e0	<,J	P	3.4e0	<	l 'n	5.2e0	<,J	5.6e0	5.2e0		P	4.3e0	3.5e0
bis(2-Ethylhexyl)phthalate	117-81-7	3.8e1	<,J	A	7.2e1	<	P	4.2e1	<,J	A	4.9e1	<,J	Ä	7.2e1	<	8.1e1	7.2e1		P	4.0e1	6.1e1
4-Bromophenyl-phenylether	101-55-3	3.0e0	<	N N	3.1e0	<	N	5.8e0	<,J	P	2.9e0	<	N N	5.8e0	<,J	6.5e0	5.8e0	- +	P	4.4e0	3.0e0
Butylbenzylphthalate	85-68-7	4.2e0	<	l n	4.3e0	<	l ii	6.1e0	<,J	P	4.0e0	<	N N	6.1e0	<,J	6.6e0	6.1e0	l	P	5.1e0	4.1e0
Carbazole	86-74-8	4.2e0	<	l n l	4.3e0	<	N I	5.8e0	<,J	P	4.0e0	<	N	5.8e0	<,J	6.2e0	5.8e0		P	5.0e0	4.1e0
4-Chloro-3-methylphenol	59-50-7	5.4e0	<	N	5.6e0	<	N	1.0e1	<,J	N	5.2e0	<	N N	1.0e1	-,∪ <,J	1.1e1	1.0e1	J	N	7.7e0	5.4e0
4-Chloroaniline	106-47-8	2.9e1	<	N	2.9e1	<del>-</del> -	N	3.6e1	<	N	2.8e1	<del>`</del>	N	3.6e1	<	3.7e1	3.6e1	-+	N	3.2e1	2.9e1
2-Chloronaphthalene	91-58-7	3.0e0	<	N	3.0e0	<	N	4.8e0	<,J	P	2.9e0	<	N	4.8e0	<,J	5.3e0	4.8e0	J	P	3.9e0	3.0e0
2-Chlorophenol	95-57-8	3.5e0	<	N	3.6e0	<	l 'n	4.8e0	<,J	P	3.4e0	<	N	4.8e0	<,J	5.2e0	4.8e0	J	P I	4.2e0	3.5e0
4-Chlorophenyl phenyl ether	7005-72-36	3.5e0	<	l 'n l	3.6e0	<	N N	5.5e0	<,J	P	3.4e0	< .	N	5.5e0	-,∪ <,J	6.0e0	5.5e0	- 1	P	4.5e0	3.5e0
Chrysene	218-01-9	3.8e0	<	N	3.9e0	<del>-</del>	N	6.8e0	<,J	P	3.7e0	<	N	6.8e0	<,J	7.5e0	6.8e0		Ė	5.3e0	3.8e0
Di-n-butylphthalate	84-74-2	3.2e1	<,J	P	3.3e1	<,J	P	3.6e1	۰,۵ ار,>	P	3.1e1	<,J	P	3.6e1	<,J	3.7e1	3.6e1	J	P	3.4e1	3.2e1
Di-n-octylphthalate	117-84-0	5.1e1	<,J	P	5.2e1	<,J	P	5.2e1	<,J	l p	4.9e1	<,J	P	5.2e1	<,J	5.4e1	5.2e1	1	P	5.1e1	5.1e1
Dibenz(a,h)anthracene	53-70-3	5.4e1	<	N	5.6e1	<	N	5.5e1	<,J	l P	5.2e1	<	N	5.6e1	<	5.7e1	5.6e1	- 1	P	5.5e1	5.4e1
Dibenzofuran	132-64-9	3.5e0	<	N	3.6e0	<	N	5.5e0	<,J	Р	3.4e0	<	N	5.5e0	<,J	6.0e0	5.5e0		Р	4.5e0	3.5e0
1,2-Dichlorobenzene	95-50-1	3.5e0	<	N	3.6e0	<	N	5.2e0	<,J	Р	3.4e0	<	N	5.2e0	<,J	5.6e0	5.2e0		Р	4.3e0	3.5e0
1,3-Dichlorobenzene	541-73-1	3.8e0	<	N	3.9e0	<	N	5.5e0	<, J	P	3.7e0	<	N	5.5e0	<,J	5.9e0	5.5e0		Р	4.7e0	3.8e0
1,4-Dichlorobenzene	106-46-7	5.1e0	<,J	P	3.9e0	<,J	P	5.2e0	ل,>	P	6.5e0	ل,>	Р	6.5e0	<,J	7.2e0	6.5e0		Р	5.1e0	5.2e0
3,3'-Dichlorobenzidine	91-94-1	3.5e1	<	N	3.6e1	<	N	4.2e1	<	N	3.4e1	<	N	4.2e1	<	4.4e1	4.2e1		N	3.9e1	3.5e1
2,4-Dichlorophenol	120-83-2	4.2e0	<	N	4.3e0	<	N	5.8e0	<	N	4.0e0	<	N	5.8e0	<	6.2e0	5.8e0	- 1	N	5.0e0	4.1e0
Diethylphthalate	84-66-2	5.1e0	۷,>	P	4.9e0	<	N	7.1e0	<,J	Р	4.6e0	<	N	7.1e0	<,J	7.7e0	7.1e0	- 1	Р	6.1e0	4.8e0
Dimethyl phthalate	131-11-3	3.1e0	<	N	3.1e0	<	N	5.2e0	<,J	Р	3.0e0	<	N	5.2e0	<,J	5.7e0	5.2e0		Р	4.1e0	3.0e0
2,4-Dimethylphenol	105-67-9	1.9e1	<	N	1.9e1	<	N	2.0e1	<	N	1.8e1	<	N	2.0e1	<	2.1e1	2.0e1		N	1.9e1	1.9e1
4,6-Dinitro-2-methylphenol	534-52-1	4.2e1	<	N	4.3e1	<	N	4.5e1	<	N	4.0e1	<	N	4.5e1	<	4.7e1	4.5e1		N	4.3e1	4.1e1
2,4-Dinitrophenol	51-28-5	8.6e1	<	N	8.8e1	<	N	9.0e1	<	N	8.3e1	<	N	9.0e1	<	9.3e1	9.0e1		N	8.8e1	8.6e1
2,4-Dinitrotoluene	121-14-2	4.2e0	<	N	4.3e0	<	N	6.1e0	<,J	P	4.0e0	<	N	6.1e0	<,J	6.6e0	6.1e0		P	5.1e0	4.1e0
2,6-Dinitrotoluene	606-20-2	3.8e0	<	N	3.9e0	<	N	5.2e0	<,J	Р	3.7e0	<	N	5.2e0	<,J	5.5e0	5.2e0		P	4.5e0	3.8e0
1,2-Diphenylhydrazine	122-66-7	3.1e0	<	N	3.2e0	<	N i	5.2e0	<,J	P	3.0e0	<	N	5.2e0	<,J	5.7e0	5.2e0	1	P	4.2e0	3.1e0
Fluoranthene	206-44-0	3.2e0	<	N I	3.3e0	<	N	5.5e0	<,J	P	3.1e0	<	N	5.5e0	<,J	6.1e0	5.5e0	- 1	P	4.3e0	3.2e0
Fluorene	86-73-7	3.2e0	<	N N	3.3e0	<	N.	5.2e0	<,J	P	3.1e0	<	N N	5.2e0	<,J	5.7e0	5.2e0		P P	4.2e0	3.2e0
Hexachlorocyclopentadiene	77-47-4	5.1e1	<	N I	5.2e1	< <	N	5.8e1	<	P P	4.9e1	<	N	5.8e1	<	6.0e1	5.8e1		P	5.5e1	5.1e1
Hexachlorobenzene Hexachlorobutadiene	118-74-1 87-68-3	3.5e0 4.8e0	< <	N N	3.6e0 4.9e0	`	N N	5.2e0	<,J	P	3.4e0	< <	N	5.2e0	<,J	5.6e0	5.2e0 6.5e0		P	4.3e0 5.6e0	3.5e0 4.8e0
Hexachloroethane	67-72-1	5.1e0	~	N	5.2e0	`	N	6.5e0 6.5e0	<,J	F	4.6e0 4.9e0	`	N	6.5e0 6.5e0	<,J	6.9e0 6.8e0	6.5e0		- F	5.8e0	5.1e0
Indeno(1,2,3-cd)pyrene	193-39-5	4.8e1	~	N	4.9e1	~	N	4.8e1	<,J <,J	P	4.6e1	~	N	4.9e1	<,J <	5.0e1	4.9e1	-	- <del>-</del> -	4.8e1	4.8e1
Isophorone	78-59-1	3.2e0	<	N	3.3e0	<	N	5.2e0	-,5 <,J	'p	3.1e0	<	N	5.2e0	<,J	5.7e0	5.2e0		' <sub>-</sub>	4.2e0	3.2e0
2-Methylnaphthalene	91-57-6	3.2e0	<	N	3.3e0	<	N	4.8e0	-,u <,J	'p	3.1e0	<	N	4.8e0	<,J	5.3e0	4.8e0		P	4.0e0	3.2e0
2-Methylphenol	95-48-7	1.5e1	<	N	1.5e1	ζ	N	1.6e1	<	N	1.4e1	~	Ň	1.6e1	<,0	1.7e1	1.6e1		N	1.6e1	1.5e1
3-Methylphenol & 4-Methylphenol	65794-96-9	1.1e1	<	N	1.1e1	<	N	1.3e1	<	N	1.0e1	<	N	1.3e1	<	1.3e1	1.3e1		N N	1.2e1	1.1e1
N-Nitroso-di-n-propylamine	621-64-7	3.5e0	<	N	3.6e0	<	N	4.8e0	<	N	3.4e0	<	N	4.8e0	<	5.2e0	4.8e0		N	4.2e0	3.5e0
N-Nitrosodimethylamine	62-75-9	3.5e0	<	N	3.6e0	<	N	4.8e0	<,J	P	3.4e0	<	N	4.8e0	<,J	5.2e0	4.8e0		Р	4.2e0	3.5e0
N-Nitrosodiphenylamine	86-30-6	4.8e0	<	N	4.9e0	<	N	6.8e0	<,J	P	4.6e0	<	N	6.8e0	<,j	7.3e0	6.8e0		P	5.8e0	4.8e0
Naphthalene	91-20-3	3.5e0	<	N	3.6e0	<	N	5.5e0	<,J	P	3.4e0	<	N	5.5e0	<,J	6.0e0	5.5e0		P	4.5e0	3.5e0
2-Nitroaniline	88-74-4	3.5e0	<	N	3.6e0	<	N	5.5e0	<	N	3.4e0	<	N	5.5e0	<	6.0e0	5.5e0	- [	N	4.5e0	3.5e0
3-Nitroaniline	99-09-2	1.3e1	<	N	1.3e1	<	N	1.6e1	<	N	1.3e1	<	N	1.6e1	<	1.7e1	1.6e1		N	1.5e1	1.3e1
4-Nitroaniline	100-01-6	1.2e1	<	N	1.2e1	<	N	1.4e1	<	N	1.1e1	<	N	1.4e1	<	1.5e1	1.4e1		N	1.3e1	1.1e1
Nitrobenzene	98-95-1	5.1e0	ار,>	Р	4.9e0	<,J	Р	6.5e0	<,J	P	3.7e0	<,J	Р	6.5e0	<,J	7.3e0	6.5e0	1	Р	5.8e0	4.3e0
2-Nitrophenol	88-75-5	2.9e1	<	P	1.3e1	<,J	Р	1.2e1	<,J	P	1.6e1	<,J	P	2.9e1	<	3.3e1	2.9e1	1	Р	2.1e1	1.5e1
4-Nitrophenol	100-02-7	2.4e1	٧,>	P	1.8e1	<	N	2.0e1	<	N	1.7e1	<,J	P	2.4e1	<,J	2.6e1	2.4e1	•	Р	2.2e1	1.7e1
2,2'-Oxybis(1-chloropropane)	108-60-1	4.5e0	<	N	4.6e0	<	N	7.1e0	<,J	Р	4.3e0	<	N	7.1e0	<,J	7.8e0	7.1e0		Р	5.8e0	4.4e0
Pentachlorobenzene	608-93-5	3.2e0	<	N I	3.3e0	<	N	4.8e0	<	N	3.1e0	<	N	4.8e0	<	5.3e0	4.8e0		N ti	4.0e0	3.2e0
Pentachloronitrobenzene	82-68-8	3.5e0	<	N I	3.6e0	<	N	5.2e0	<	N	3.4e0	<	N	5.2e0	<	5.6e0	5.2e0		N	4.3e0	3.5e0
Pentachlorophenol	87-86-5	9.6e1	<	N	9.8e1	<	N I	1.0e2	<	N	1.1e2	<	N	1.1e2	<	1.2e2	1.1e2	ŀ	N	9.8e1	1.1e2
Phenanthrene	85-01-8	3.1e0	<	N	3.2e0	< .	<u>N</u>	5.5e0	<,J	P	3.0e0	< .	N	5.5e0	ال,>	6.1e0	5.5e0		P	4.3e0	3.1e0
Phenol	108-95-2	2.5e1	<	P	1.2e1	<,J	P	1.1e1	<,J	Р	1.2e1	<,J	P	2.5e1	<	2.8e1	2.5e1		Р	1.8e1	1.2e1
Pyrene	129-00-0	3.2e0	<	N	3.3e0	<	N	5.8e0	<,J	P	3.1e0	<b> </b>	N	5.8e0	<,J	6.5e0	5.8e0		P	4.5e0	3.2e0
Pyridine	110-86-1	5.4e0	<	N	5.6e0	< <	N I	9.0e0	<u> </u>	N	5.2e0	<	N	9.0e0	<	1.0e1	9.0e0	- 1	N	7.2e0	5.4e0
1,2,4,5-Tetrachlorobenzene	95-94-3	3.5e0	<	N N	3.6e0	<	N N	5.2e0	<	N P	3.4e0	<	N N	5.2e0	<	5.6e0	5.2e0	<b> </b> -	N P	4.3e0	3.5e0 3.8e0
1,2,4-Trichlorobenzene 2,4,5-Trichlorophenol	120-82-1 95-95-4	3.8e0	< <	N N	3.9e0 8.2e0	`	N N	5.5e0	<,J <		3.7e0	· _	N N	5.5e0	<,J <	5.9e0	5.5e0	1	N	4.7e0 8.7e0	7.9e0
2,4,5-1 nchlorophenol 2,4,6-Trichlorophenol	95-95-4 88-06-2	8.0e0 5.1e0	< <	N N	5.2e0 5.2e0	~	N	9.4e0 6.8e0	<b>~</b>	N N	7.7e0 4.9e0	<	N	9.4e0 6.8e0	<b>~</b>	9.8e0 7.2e0	9.4e0 6.8e0		N I	5.9e0	7.9e0 5.1e0
z,+,o-monorophenor	00-00-2	5. TeU		14	3.2 <del>0</del> 0	`	iN.	0.080	`	14	4.5€∪	`	IN	0.000	`	7.2e0	0.860		14	5.960	3.160

Table B-22. SVOC concentration-basis

TICs	CAS Registry Number	STRT-1		Project Specific Flag	END-1		Project Specific Flag	STRT-2 µg/dscm		Project Specific Flag	END-2 µg/dscm		Project Specific Flag	Max value µgidsom		Aug+Ze µg/dsom	Results µg/tsom	Project Specific Flag	STRT Hure Avgs ug/dscm	Avgs.
1,3-Bercenedicarboxylic acid, 2,4-Hexadiene 2,5-Diethylphenol 2,Cyclohoxene-1-one, 3-methyl-	1330-96-7 592-46-1 876-20-0 1193-18-6				5.2e1	NJM	P	2.7e0	N.J.Q	P	8.6e0 3.5e1 9.8e-1	MTN WTN	p. p. a	2.7e0 5.2e1 3.1e1 9.8e-1	MT'N MT'N MT'N	9.2e1	2.7e0 5.2e1 3.1e1 9.8e-1	P P P	2.7e0	3.1e1 3.1e1 9.8e-1
2-hexanore 3-Hexanore Beroskitchyde Beroskitchyde, 4-ethyl-	591-78-6 589-30-8 100-52-7 4748-78-1	7.4e1 6.1e1 3.5e2	M,LM M,LM M,LM	P P	3.9e1 3.1e1 2.4e2 1.9e1	M,L,M M,L,M M,L,M M,L,M	200	3.1e1 2.2e2	NJ,M NJ,M	P	2.2e1 2.2e2	N.I.M M.I.M	P	7,4e1 6,1e1 3,5e2 1,9e1	MLN MLN MLN	1.0e2 7.0e1 3.6e2	7,4e1 6.1e1 3.5e2 1.9e1	PPP	7.4e1 4.6e1 2.8e2	3.9e1 2.6e1 2.3e2 1.9e1
Berozidehyde, ethyl- Berozojejpyrene Cyclododecane Dodecane	53951-50-1 192-97-2 294-62-2 112-40-3	9.9e0 2.1e1	M,LM M,LM	P	9.8e0 1.6e1	NJ,M NJ,M	p	4.5e0 1.8e0 1.8e1	N.Y.W N.Y.O	0.00	1.6e1 4.0e0 1.1e1	N,I,M N,I,M M,I,M	P	1.9e1 4.5e0 9.9e0 2.1e1	ML/A ML/A ML/A	1.5e1 2.5e1	1.9e1 4.5e0 9.9e0 2.1e1	0 0 0 0	4.5e0 5.9e0 2.0e1	1.9e1 6.9e0 1.4e1
Elcosane Formic acid, phenylmethyl ester Furan, 2,5-dimethyl- Henelcosane	112-95-8 104-57-4 625-86-5 629-94-7	2.0e0 4.5e0 6.4e-1	N.J.M N.J.M	0 0 0	2.4e1 2.6e0	NJM	P	3.2e0	N,J,M	P	5.6e0 2.9e1 2.8e0	MLN MLN	P	5.8e0 2.9e1 4.5e0 2.6e0	MLW MLW MLW MLW	9.4e0 3.4e1 5.4e0 4.8e0	5.8e0 2.9e1 4.5e0 2.8e0	200	2.0e0 3.8e0 6.4e-1	5.5e0 2.6e1 2.6e0 2.8e0
Heptacosane Heptadocane Heptane, 2,3-dimethyl- Heptane, 2,5-dimethyl-	593-49-7 629-78-7 3074-71-3 2216-30-0	2.8e0	M,I,M	P	1.3e0 3.2e0	N.I.M N.I.M	P P	2.360	N.J.M	p	2.6e1	N,J,M	P	2.5e1 2.5e0 3.2e0 2.3e0	MLM MLM MLM MLM	4.3+0	2.6e1 2.6e0 3.2e0 2.3e0	0.0.0	2.8e0 2.3e0	2.6e1 1.3e0 3.2e0
Hesacosane Hesadecanoic acid Hesanedicic acid, bis(2-ethylh) Hesatriacontane	630-01-3 57-10-3 103-23-1 630-06-8	1.7e0	M,I,M	Р				3.2e0	N,J,Q	P	2.0e1	NJM	P	2.0e1 1.7e0 3.2e0 1.8e0	MLM DLM MLM		2.0e1 1.7e0 3.2e0 1.8e0	9 9	1.7e0 3.2e0	2.0e1
Naphthalene, 1-methyl- Octadecanoic acid Octane, 3-methyl- Pentacosane	90-12-0 57-11-4 2216-33-3 629-99-2	1.2e0 2.5e0	NJ,M	P	6.5e-1	M,L,M	P	2.5e0	D,L,M	р	1.1e1	NJM	P	2.5e0 1.2e0 2.5e0 1.1e1	MLM MLM MLM	1.6e0	2.5e0 1.2e0 2.5e0 1.1e1	p p p	2.5e0 1.2e0 2.5e0	6.5e-1 1.1e1
Pertadecane Phosphine oxide, triphenyl- Phosphoric acid tributyl ester Tetracospres	629-62-9 791-28-6 126-73-8 646-31-1	4.5e0 2.6e0 7.4e0	M'T'N M'T'N M'T'N		5.2eG	NJM	Р				7.4e0 8.0e0	N.I.M N.I.M	0.0	4.8e0 2.8e0 7.4e0 8.0e0	NJM NJM NJM	7.9e0 9.1e0	4.8e0 2.8e0 7.4e0 8.0e0	P P P	4.8e0 2.8e0 7.4e0	1.4eG 6.3eG 8.0eG
Tetradecane Tetratetracontane Tridecane	629-59-4 7096-22-8 629-50-5	2.8e0 2.1e0 5.4e0	MALM	P	3.1e1 6.9e0	N.J.M N.J.M	P	1.5e1 5.5e0	NJ,M	P	3.0e1 6.5e0	M.L.N	P	3.1e1 2.1e0 6.9e0	N,JM N,JM	4.7u1 7:3e0	3.1e1 2.1e0 6.9e0	p p	1,0e1 2,1e0 6,1e0	3.1e1 6.7e0

Table B-23. 0031 concentration-basis.

Marche March 1969   March 1969	Table B-23. 0031 concentration	CAS	STRT-1	П	Project	END-1	Ē	Project	STRT-2	71	Project	END-2	77	Project	Max value	_	Avg+2σ 🔟	Results		Project	STRIRUN	END Run
Process	Analyte	Registry Number		~~	Specific		lag	Specific		Flag	Specific		Flag	Specific		Flag	- 0)		Flag			
Accordance   17-14-1   3.60   c   N   3.90   c   N	Acetone		8.9e1	В	<del></del>	9.2e1	В		6.3e1	J,B		4.8e1	<,J,B		6.3e1	J.B	1.2e2	6.3e1	J.B			
	Acrylonitrile	107-13-1	3.5e1	<	N	3.6e1	<	N	3.5e1		N			N	1						3.5e1	3.6e1
Binney Commonwhale	Benzene	71-43-2	4.9e0	<	P	3.1e0	<	P	2.8e0	<,J	Р	2.6e0	<	P	4.9e0	<	5.5e0	4.9e0	<	Р	3.9e0	2.8e0
Binomothermethene   72,574   5,60   c   N   1,50   c   N   1,50   c   N   1,50   c   N   1,50   c   N   1,50   c   N   1,50   c   N   1,50   c   N   1,50   c   N   1,50   c   N   2,50	Bromobenzene	108-86-1	1.5e0	<	N	1.5e0	<	N	1.5e0	<	N	1.5e0	<	N	1.5e0	<	1.6e0	1.5e0	<	N I	1.5e0	1.5e0
Secondary   7-55-2   2-20   C   N	Bromochloromethane		1.9e0	<	N	1.9e0	<	N	1.9e0	<	N	1.9e0	<	N	1.9e0	<	1.9e0	1.9e0	<	N	1.9e0	1.9e0
State	Bromodichloromethane			<				N		<		1.5e0	<	N	1.5e0	<	1.6e0	1.5e0	<	N	1.5e0	1.5e0
28-bissone	B .					L		1							1	<	B .		<			
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See-Euglymanne   See-															1							
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Chlorothermore   124-841   1.900								l ' l			i '											
Chlorophomorethame					i									1	•							
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12-Disconnections	4-Chlorotoluene	106-43-4	7.3e-1	<	N	7.5e-1	<	N		<			<	N				4				
Distromomenhane	1,2-Dibromo-3-chloropropane	96-12-8	3.3e0	<	N	3.6e0	<	N	3.5e0	<	N	3.4e0	<	N	3.6e0	<	3.7e0	3.6e0	<	N	3.4e0	3.5e0
1,2-Dichroreheremen	1,2-Dibromoethane	106-93-4	2.5e0	<	N	2.6e0	<	N	2.5e0	<	N	2.6e0	<	N	2.6e0	<	2.6e0	2.6e0	<	N	2.5e0	2.6e0
1,3-Dichtoropeneme	Dibromomethane			<			<	N	2.1e0	<	N	2.2e0	<	N	2.2e0	<	2.2e0	2.2e0	<	N	2.1e0	2.2e0
1.4-Dichrorbenzemen	.,		1	<			<		2.3e0	<		2.3e0	<		2.3e0	<	2.3e0	2.3e0	<		2.2e0	2.3e0
Dichlorodifucomethane   75-71-8   32-80   <   P   2.0e0   <   P   2.5e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <	1	1										i							<			
11-Dichloremane																	1	1				
12-Dichforcethene																						
11-Dichforophene   75-35-4   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P   2.0e0   <   P																						
1986-99-2   1.9e0   x	I '														1							
Tams-1,2-Dichloroethene	1 *				1 '													B .		''		
12-Dichloropropane																						
13-Dichloropropage	•		1												•							
22-Dichloropropane		1			1			N												N		
1.1-Dichloropropen																						
clis-13-Dichloropropee         1061-01-5         1.8e0          N         1.7e0          N         1.7e0          N         1.7e0          N         1.7e0          N         1.7e0          N         1.9e0          N         2.9e0          N         1.9e0          N         2.9e0          N         2.9e0				<																		
Trans-1,3-Dichloropropene   10061-02-6   1,9e0		10061-01-5	i e	<	N		<	N		<	N		<	N		<	i e					
Hexachlorobutadiene	trans-1,3-Dichloropropene	10061-02-6	1.9e0	<	N	1.9e0	<	N	1.9e0	<	N	1.9e0	<	N		<	1.9e0		<	N	1.9e0	1.9e0
2-Hexanore 591-78-6 8.9e0 < N 7.1e0 < N 7.0e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0 < N 7.1e0	Ethylbenzene	100-41-4	1.2e0	<	N	1.2e0	<	N	1.2e0	<	N	1.2e0	<	N	1.2e0	<	1.2e0	1.2e0	<	N	1.2e0	1.2e0
Septemble   98.82-8   8,7e-1   C   N   8.9e-1   C   N   8.8e-1   C   N   8.9e-1   C   N   7.9e   C   N   7.9e   C   N   7.9e   C   N   7.9e   C   N   7.9e   C   N   7.9e   C   N   7.9e   C   N   7.9e   C	Hexachlorobutadiene	87-68-3	2.7e0	<	N	2.8e0	<	N	2.8e0	<	N	2.9e0	<	N	2.9e0	<	3.0e0	2.9e0	<	N	2.7e0	2.9e0
p-Isopropytoluene 99-87-6 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N 1.4e0 < N	2-Hexanone		6.9e0	<		7.1e0	<	N	7.0e0	<	N	7.1e0	<	N	7.1e0	<	7.3e0	7.1e0	<	N	7.0e0	7.1e0
Methylene chloride	Isopropylbenzene									<		8.9e-1	<	N	8.9e-1	<	9.1e-1	8.9e-1	<			
4-Methyl-2-pentanone		~ · · · · · · · · · · · · · · · · · · ·																				
Naphthalene 91-20-3 2.6e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0 < N 2.7e0	1 '							l 1										1				1
103-65-1   8.3e-1						_																
Styrene   100-42-5   9.7e-1   <   N   1.0e0   <   N   9.8e-1   <   N   9.9e-1   <   N   1.0e0   <   N   9.9e-1   <   N   1.0e0   <   N   9.7e-1   <   N   1.0e0   <   N   9.7e-1   <   N   1.0e0   <   N   9.7e-1   <   N   1.0e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N   1.3e0   <   N	B '										1		- 1									
1,1,1,2-Tetrachloroethane						~	_															
1,1,2,2-Tetrachloroethane														,,,							1	1
Tetrachloroethene 127-18-4 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N 1.9e0 < N																						
Toluene 108-88-3	., ., .,																					
1,2,3-Trichlorobenzene																						
1,2,4-Trichlorobenzene																						
1,1,1-Trichloroethane	1,2,4-Trichlorobenzene																					1 1
1,1,2-Trichloroethane	1,1,1-Trichloroethane			<						•												
Trichloroethene 79-01-6 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.0e0 < N 2.	1,1,2-Trichloroethane			<		··-	<			<			<			<			<			
1,2,3-Trichloropropane       96-18-4       3.0e0        N       3.1e0        N       1.3e0        N       1.3e0        N       1.3e0        N       1.3e0        N       7.5e-1        N       7.5e-1        N       7.5e-1        N       7.5e-1        N       7.5e-1        N       7.5e-1        N       7.5e	Trichloroethene	79-01-6	2.0e0	<	N	2.0e0	<	N	2.0e0	<	N		<	N		<			<	N	2.0e0	2.0e0
1,2,4-Trimethylbenzene 95-63-6 1.2e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0 < N 1.3e0	Trichlorofluoromethane	75-69-4	2.0e0	٧,>		2.0e0	<,J	Р	2.0e0	<,J	P	2.0e0	<,J	Ρ	2.0e0	<,J	2.1e0	2.0e0	<,J	Р	2.0e0	2.0e0
1,3,5-Trimethylbenzene	1,2,3-Trichloropropane		3.0e0	<		3.1e0	<		3.0e0		<u>N</u>	3.1e0	<	N	3.1e0		3.1e0	3.1e0		N	3.0e0	3.1e0
Vinyl chloride 75-01-4 1.6e0 <,J P 2.3e0 <,J P 1.6e0 <,J P 2.2e0 <,J P 2.3e0 <,J P 2.3e0 <,J P 2.3e0 <,J P 1.6e0 2.2e0   m-Xylene & p-Xylene	1,2,4-Trimethylbenzene			<					1.3e0	<	N	1.3e0	<	N	1.3e0	<	1.3e0	1.3e0	<	N	1.2e0	
m-Xylene & p-Xylene 136777-61-2 5.4e0 < N 5.6e0 < N 5.5e0 < N 5.6e0 < N 5.6e0 < 5.7e0 5.6e0 < N 5.5e0 5.6e0	1,3,5-Trimethylbenzene									- 1									<			
	Vinyl chloride							-			- 1			,								
o-xyiene 95-47-6 9.5e-1 < N 1.0e0 <,J P 9.8e-1 < N 9.8e-1 < J P 1.0e0 <,J 1.0e0   1.0e0 <,J P 9.6e-1 9.9e-1	m-Xylene & p-Xylene																					
	o-Xylene	95-47-6	9.5e-1	<	N	1.0e0	<,J	Р	9.8e-1	<	N	9.8e-1	<,J	Р	1.0e0	<,J	1.0e0	1.0e0	<,J	Р	9.6e-1	9.9e-1

Table B-23. 0031 concentration-basis.

TICs	CAS Registry Number	STRT-1 µg/dscm		Project Specific Flag	END-1 μg/dscm	Flag	Project Specific Flag	STRT-2 µg/dscm	Flag	Project Specific Flag	END-2 μg/dscm	Flag	Project Specific Flag	Max value μg/dscm	Flag	Avg+2σ μg/dscm	Flag	Results μg/dscm	Flag	Project Specific Flag	STRT Run Avgs µg/dscm	END Run Avgs µg/dscm
1-Heptene	592-76-7	6.7e-1	N,J,M	Р											N,J,M			6.7e-1	N,J,M	Р	6.7e-1	
Benzonitrile	100-47-0				2.4e0	N,J,M	Р	9.0e-1	N,J,M	P	6.0e-1	N,J,M	Р	2.4e0	N,J,M	3.3e0	- 1	2.4e0	N,J,M	Р	9.0e-1	1.5e0
Butane, 1-chloro-	109-69-3	7.1e-1	N,J,M	P	l									7.1e-1	N,J,M		- 1	7.1e-1	N,J,M	Р	7.1e-1	
Cyclobutane, ethenyl-	2597-49-1	L			1.5e0	N,J,M	P							1.5e0	N,J,M			1.5e0	N,J,M	P	1 1	1.5e0
Cyclohexane, hexyl-	4292-75-5	7.3e-1	N,J,M	Р										7.3e-1	N,J,M			7.3e-1	N,J,M	Р	7.3e-1	
Cyclohexane, methyl-	108-87-2	1.4e0	N,J,M	P	2.6e0	N,J,M	Р	2.5e0	N,J,M	Р	1.5e0	N,J,M	Р	2.6e0	N,J,M	3.2e0		2.6e0	N,J,M	Р	1.9e0	2.0e0
Cyclohexene	110-83-8	1.2e0	N,J,M	Р	4.6e-1	N,J,M	Р	2.3e0	N,J,M	P	3.3e-1	N,J,M	Р	2.3e0	N,J,M	2.8e0		2.3e0	N,J,M	Р	1.7e0	4.0e-1
Cyclopentane, 1,2-dimethyl-	2452-99-5				l			6.6e-1	N,J,M	P				6.6e-1	N,J,M	9.9 <del>e</del> -1		6.6e-1	N,J,M	Р	3.3e-1	1 1
Cyclopentane, 1,2-dimethyl-, t	822-50-4			i	1.0e0	N,J,M	Р				3.8e-1	N,J,M	Р	1.0e0	N,J,M	1.6e0		1.0e0	N,J,M	Р		7.0e-1
Cyclopentane, ethyl-	1640-89-7	3.5e-1	N,J,M	P	5.2e-1	N,J,M	Р	4.5e-1	N,J,M	Р				5.2e-1	N,J,M	7.9e-1		5.2e-1	N,J,M	Р	4.0e-1	2.6e-1
Decane	124-18-5	1.5e0	N,J,M	P										1.5e0	N,J,M			1.5e0	N,J,M	Р	1.5e0	1 1
Decane, 2,2,5-trimethyl-	62237-96-1							7.5e-1	N,J,M	Р				7.5e-1	N,J,M			7.5e-1	N,J,M	Р	7.5e-1	1 1
Decane, 2,9-dimethyl-	1002-17-1	7.9e-1	N,J,M	Р										7.9e-1	N,J,M			7.9e-1	N,J,M	Р	7.9e-1	
Dodecane	112-40-3	3.8e2	N,J,M	P	3.3e2	N,J,M	Р	2.8e2	N,J,M	Р	2.2e2	N,J,M	Р		N,J,M	4.5e2		3.8e2	N,J,M	Р	3.3e2	2.7e2
Dodecane, 6-methyl-	6044-71-9							6.5e-1	N,J,M	Р					N,J,M			6.5e-1	N,J,M	Р	6.5e-1	
Hexadecane	544-76-3				2.7e0	N,J,M	Р							2.7e0	N,J,M			2.7e0	N,J,M	P	1 1	2.7e0
Hexane, 2,4-dimethyl-	589-43-5	1.4e0	N,J,M	Р	2.3e0	N,J,M	Р	2.5e0	N,J,M	Р	1.2e0	N,J,M	Р	2.5e0	N,J,M	3.2e0		2.5e0	N,J,M	Р	1.9e0	1.7e0
Hexane, 2-methyl-	591-76-4	2.1e0	N,J,M	P	4.0e0	N,J,M	Р	3.8e0	N,J,M	Р	3.1e0	N,J,M	Р	4.0e0	N,J,M	4.9e0	- 1	4.0e0	N,J,M	Р	2.9e0	3.5e0
Hexane, 3-methyl-	589-34-4	4.7e0	N,J,M	Р	9.7e0	N,J,M	Р	3.4e0	N,J,M	Р	4.3e0	N,J,M	Р	9.7e0	N,J,M	1.1e1		9.7e0	N,J,M	Р	4.0e0	7.0e0
Methane, trichloronitro-	76-06-2	į			4.6e0	N,J,M	Р							4.6e0	N,J,M			4.6e0	N,J,M	Р	1 1	4.6e0
Octane	111-65-9	3.3e-1	N,J,M	Р											N.J.M			3.3e-1	N,J,M	Р	3.3e-1	
Pentane, 2,3-dimethyl-	565-59-3	2.2e0	N,J,M	Р	4.0e-1	N,J,M	Р	1.4e0	N,J,M	Р	1.8e0	N,J,M	Р	2.2e0	N,J,M	3.0e0	1	2.2e0	N,J,M	Р	1.8e0	1.1e0
Pentane, 3,3-dimethyl-	562-49-2				7.5e-1	N,J,M	Р								N.J.M	9.4e-1	ı	7.5e-1	N,J,M	Р		3.8e-1
Pentane, 3-ethyl-	617-78-7							4.3e-1	N,J,M	Р	4.1e-1	N.J.M	Р	4.3e-1	N,J,M	6.9e-1	I	4.3e-1	N,J,M	Р	2.1e-1	2.0e-1
Tetradecane	629-59-4	1.4e0	N,J,M	Р	1.3e1	N,J,M	P	1.2e1	N,J,M	Р	1.8e1	N,J,M	P		N.J.M	2.5e1		1.8e1	N,J,M	Р	6.8e0	1.5e1
Tridecane	629-50-5	3.5e1	N,J,M	Р	4.0e1	N,J,M	Р	1.0e0	N,J,M	P	4.0e1	N,J,M	P		N,J,M	6.6e1		4.0e1	N,J,M	Р	1.8e1	4.0e1
Undecane	1120-21-4	1.2e1	N,J,M	Р	6.4e0	N,J,M	Р	3.0e0	N,J,M	Р	1.5e0	N,J,M	P		N,J,M	1.5e1		1.2e1	N J M	P	7.7e0	4.0e0
Undecane, 2,6-dimethyl-	17301-23-4	1.4e0	N,J,M	Р				5.8e-1	N,J,M	Р	1.1e0	N,J,M	P		N,J,M	1.8e0		1.4e0	N,J,M	P	9.7e-1	1.1e0
Undecane, 5-methyl-	1632-70-8	7.9e0	N,J,M	Р	2.3e0	N,J,M	P	2.6e0	N,J,M	Р	2.3e0	N,J,M	P		N,J,M	9.3e0		7.9e0	N,J,M	Р	5.3e0	2.3e0

Table B-24. 0050 concentration-basis.

Analyte	STRT-1 ppmv, dry	Flag	END-1 ppmv, dry	Flag	STRT-2 ppmv, dry	Flag	END-2 ppmv, dry	0)	Max value ppmv, dry	Flag	Avg+2o ፲ ppmv, dry 🚨	Results ppmv, dry	Flag	STRT Run Avgs ppmv, dry	END Run Avgs ppmv, dry
Chloride (as HCl) Chloride (as Cl2)	6.1e-1 3.1e-2	B <	5.7e-1 5.7e-2	B B	6.4e-1 3.2e-2	B <	6.2e-1 3.2e-2	B <	6.4e-1 5.7e-2	B B	6.7e-1 6.4e-2	6.4e-1 5.7e-2	B B	6.3e-1 3.2e-2	6.0e-1 4.5e-2
Particulate	mg/dscm 1.1e0	В	mg/dscm 8.6e-1	В	mg/dscm 3.7e-1		mg/dscm 2.6e-1	В	mg/dscm 1.1e0	В	mg/dscm 1.5e0	mg/dscm 1.1e0	В	mg/dscm 7.4e-1	mg/dscm 5.6e-1

Table B-25, 0060 concentration-basis.

Analyte	CAS Registry Number	STRT-1 µg/dscm	Flag	Project Specific Flags	END-1 µg/dscm	Flag	Project Specific Flags	STRT-2 µg/dscm	Flag	Project Specific Flags	END-2 µg/dscm	Flag	Project Specific Flags	Max value µg/dscm	~~	Avg+2o ਜ਼ੁ µg/dscm o	Results µg/dscm	Flag	Project Specific Flag	STRT Run Avgs µg/dscm	END Run Avgs μg/dscm
Aluminum (AI)	7429-90-5	4.6e1		Α	7.4e1		Α	2.3e1		Α	2.4e1		Α	7.4e1		9.0e1	7.4e1		Α	3.4e1	4.9e1
Antimony (Sb)	7440-36-0	1.5e0	В	A	1.4e0	В	Α	1.4e0	В	Α	1.3e0	В	Α	1.5e0	В	1.5e0	1.5e0		Α	1.4e0	1.4e0
Arsenic (As)	7440-38-2	5.2e-1	<,B	Р	4.4e-1	<	N	4.9e-1	<	N	5.2e-1	<	N	5.2e-1	<,B	5.7e-1	5.2e-1		Р	5.1e-1	4.8e-1
Barium (Ba)	7440-39-3	2.2e0	В	A	2.7e0	В	Α	1.6e0	В	A	1.5e0	В	Α	2.7e0	В	3.2e0	2.7e0		Α	1.9e0	2.1e0
Beryllium (Be)	7440-41-7	2.1e-1	<,B	Р	1.8e-1	<,B	Р	2.0e-1	<,B	Р	2.1e-1	<,B	P	2.1e-1	<,B	2.3e-1	2.1e-1		Р	2.1e-1	2.0e-1
Cadmium (Cd)	7440-43-9	1.5e-1	В	A	2.2e-1	B	Α .	8.2e-2	<,B	P	8.8e-2	<	N	2.2e-1	В	2.7e-1	2.2e-1		Р	1.2e-1	1.6e-1
Chromium (Cr)	7440-47-3	9.8e-1		A	1.3e0		Α	7.6e-1		A	1.7e0		Α	1.7e0		2.0e0	1.7e0		Α	8.7e-1	1.5e0
Cobalt (Co)	7440-48-4	8.8e-1	В	A	7.4e-1	<	N .	8.2e-1	<	N	8.8e-1	<	N	8.8e-1	В	9.7e-1	8.8e-1		P	8.5e-1	8.1e-1
Copper (Cu)	7440-50-8	1.4e0	В	Α	9.9e-1	В	A	7.0e-1	В	Α	4.6e-1	<,B	Р	1.4e0	В	1.7e0	1.4e0		Р	1.1e0	7.2e-1
Lead (Pb)	7439-92-1	4.3e-1	<,B	P	3,6e-1	В	Α	4.3e-1	<,B	P	3.9e-1	<,B	Р	4.3e-1	<,B	4.7e-1	4.3e-1		Р	4.3e-1	3.7e-1
Manganese (Mn)	7439-96-5	6.2e0		Α	6.3e0		Α	1.2e1		A	2.4e1		Α .	2.4e1		2.8e1	2.4e1		Α	8.9e0	1.5e1
Mercury (Hg)	7439-97-6	3.3e1		Α	4.1e1		Р	3.4e1		P	3.6e1		Р	4.1e1		4.3e1	4.1e1		P	3.3e1	3.9e1
Nickel (Ni)	7440-02-0	1.6e0	В	Α	1.6e0	В	Ā	1.4e0	В	Α	1.3e0	В	Α	1.6e0	В	1.8e0	1.6e0		Α	1.5e0	1.5e0
Selenium (Se)	7782-49-2	1.0e0	В	A	5.7e-1	<	Р	7.9e-1	В	A .	7.5e-1	В	Α	1.0e0	В	1.2e0	1.0e0		Р	9.2e-1	6.6e-1
Silver (Ag)	7440-22-4	8.5e-1	<	N	7.1e-1	<	N	7.9e-1	<	N j	8.5e-1	<	N	8.5e-1	<	9.3e-1	8.5e-1		N	8.2e-1	7.8e-1
Thallium (TI)	7440-28-0	8.2e-1	<	N	6.6e-1	<	N	7.6e-1	<	N	7.8e-1	· <	N	8.2e-1	<	8.9e-1	8.2e-1		N	7.9e-1	7.2e-1
Vanadium (V)	7440-62-2	8.8e-1	<	N	7.4e-1	<	N	8.2e-1	<	N	8.8e-1	<	N	8.8e-1	<	9.7e-1	8.8e-1		N	8.5e-1	8.1e-1
Zinc (Zn)	7440-66-6	2.0e1		Α	3,3e1		Α	9.1e0		A	5.6e0		Α	3.3e1		4.1e1	3.3e1		Α	1.5e1	1.9e1

 Total Metals
 1.9e2

 Total Detected Metals
 1.8e2

Table B-26. 0060 blank corrected concentration-basis.

Analyte	CAS Registry Number	STRT-1 µg/dscm	Flag	Project Specific Flags	END-1 µg/dscm	Flag	Project Specific Flags	STRT-2 µg/dscm	Flag	Project Specific Flags	END-2 μg/dscm	Flag	Project Specific Flags	Max value µg/dscm	Flag	Avg+2σ μg/dscm	Flag	Results µg/dscm	Flag	Project Specific Flag	STRT Run Avgs µg/dscm	END Run Avgs μg/dscm
Aluminum (AI)	7429-90-5	4.2e1		Α	7.0e1		Α	2.1e1		Α	2.2e1		Α	7.0e1		8,5e1		7.0e1		Α	3.2e1	4.6e1
Antimony (Sb)	7440-36-0	3.4e-1	В	Α	4.2e-1	В	A	2.9e-1	В	Α	1.8e-1	В	Α	4.2e-1	В	5.1e-1	l	4.2e-1		Α	3.1e-1	3.0e-1
Arsenic (As)	7440-38-2	2.8e-1	<,B	P	2.4e-1	<	N	2.6e-1	<	N	2.8e-1	<	N	2.8e-1	<,B	3.1e-1		2.8e-1		Р	2.7e-1	2.6e-1
Barium (Ba)	7440-39-3	7.0e-1	В	Α	1.5e0	В	Α	1.6e-1	В	Α	4.6e-2	В	Α	1.5e0	В	1.9e0		1.5e0		Α	4.3e-1	7.8e-1
Beryllium (Be)	7440-41-7	1.5e-1	<,B	Р	1.2e-1	<,B	Р	1.4e-1	<,B	P	1.5e-1	<,B	Р	1.5e-1	<,B	1.6e-1		1.5e-1		P	1.4e-1	1.4e-1
Cadmium (Cd)	7440-43-9	1.3e-1	В	Α	2.1e-1	В	Α	6. <b>4e-</b> 2	<,B	P	6.9e-2	<	N	2.1e-1	В	2.5e-1		2.1e-1		Р	9.8e-2	1.4e-1
Chromium (Cr)	7440-47-3	3.8e-1		Α .	7.9e-1		Α	2.0e-1		Α	1.1e0		Α	1.1e0		1.4e0		1.1e0		Α	2.9e-1	9.3e-1
Cobalt (Co)	7440-48-4	8.8e-1	В	Α	7.4e-1	<	N	8.2e-1	<	N	8.8e-1	<	N	8.8e-1	В	9.7e-1		8.8e-1		P	8.5e-1	8.1e-1
Copper (Cu)	7440-50-8	1.4e0	В	A .	9.9e-1	В	Α	6.9e-1	В	Α	4.5e-1	<,B	Р	1.4e0	В	1.7e0		1.4e0		Р	1.1e0	7.2e-1
Lead (Pb)	7439-92-1	2.8e-1	<,B	P	2.4e-1	В	Α .	3.0e-1	<,B	Р	2.5e-1	<,B	P	3.0e-1	<,B	3.2e-1		3.0e-1		P	2.9e-1	2.4e-1
Manganese (Mn)	7439-96-5	5.7e0		Α .	6.0 <b>e0</b>		Α .	1.1e1		Α	2.3e1		Α	2.3e1		2.7e1		2.3e1		A	8.4e0	1.4e1
Mercury (Hg)	7439-97-6	3.3e1		Α	4.1e1		Р	3.4e1		Р	3.6e1		P	4.1e1		4.3e1		4.1e1		P	3.3e1	3.9e1
Nickel (Ni)	7440-02-0	5.0e-1	В	Α	7.0e-1	В	Α	3.2e-1	В	Α	2.1e-1	В	Α	7.0e-1	В	8.6e-1		7.0e-1		Α	4.1e-1	4.5e-1
Selenium (Se)	7782-49-2	3.3e-11	В	A	2.7e-11	<	P	3.0e-11	В	Α	3.3e-11	В	A	3.3e-11	В	3.6e-11		3.3e-11		Р	3.2e-11	3.0e-11
Silver (Ag)	7440-22-4	8.5e-1	<	N	7.1e-1	<	N	8.0e-1	<	N	8.5e-1	<	N	8.5e-1	<	9.4e-1		8.5e-1		N	8.2e-1	7.8e-1
Thallium (TI)	7440-28-0	8.2e-1	<	N	6.6e-1	<	N	7.6e-1	<	N	7.8e-1	<	N	8.2e-1	<	8.9e-1		8.2e-1		N	7.9e-1	7.2e-1
Vanadium (V)	7440-62-2	8.8e-1	<	N	7.4e-1	<	N	8.2e-1	<	N	8.8e-1	<	N	8.8e-1	<	9.7e-1		8.8e-1		N	8.5e-1	8.1e-1
Zinc (Zn)	7440-66-6	1.8e1		Α	3.1e1		Α	7.9e0		Α	4.1e0		A	3.1e1		4.0e1		3.1e1		Α	1.3e1	1.8e1
Total Metals																-		1.7e2				<del></del>

Total Detected Metals

1.7e2

# APPENDIX C

PROCESS STREAM SAMPLING DATA

		,		

Table C-1. VOC target analyte list reported by the INTEC ALD.

CAS#	Compound	CAS#	Compound
74-87-3	Chloromethane	79-01 <b>-</b> 6	Trichloroethene
75-01-04	Vinyl Chloride	78-87 <b>-</b> 5	1,2-Dichloropropane
74-83-9	Bromomethane	75-27-4	Bromodichloromethane
75-00-3	Chloroethane	10061-01-5	Cis-1,3-dichloropropene
75-69-4	Trichlorofluoromethane	108-10-1	4-Methyl-2-pentanone
75-35-4	1,1-Dichloroethene	108-88-3	Toluene
76-13-1	1,1,2-Trichloro-1,2,2-	10061-02-6	Trans-1,3-dichloropropene
	trifluoroethane		
75-15-0	Carbon disulfide	79-00-5	1,1,2-Trichloroethane
67-64-1	Acetone	127-18-4	Tetrachloroethene
75-09-2	Methylene chloride	591-78-6	2-Hexanone
	(Dichloromethane)		
156-60-5	Trans-1,2-dichloroethene	124-48-1	Dibromochloromethane
75-34-3	1,1-Dichloroethane	108-90-7	Chlorobenzene
156-59-2	Cis-1,2-dichloroethene	100-41-4	Ethylbenzene
78-93-3	2-Butanone	108-38-3	M-xylene and 106-42-3 p-xylene
67-66-3	Chloroform	95-47-6	O-xylene
71-55-6	1,1,1-Trichloroethane	100-42-5	Styrene
56-23-5	Carbon tetrachloride	75-25-2	Bromoform
71-43-2	Benzene	79-34-5	1,1,2,2-Tetrachloroethane
107-06-2	1,2-Dichloroethane	· · · · · · · · · · · · · · · · · · ·	

Table C-2. SVOC target analyte list reported by the INTEC ALD.

CAS#	Compound	CAS#	Compound
62-75-9	N-Nitrosodimethylamine	83-32-9	Acenaphthene
110-86-1	Pyridine	51-28-5	2,4-Dinitrophenol
108-95-2	Phenol	100-02-7	4-Nitrophenol
111-44-4	bis(2-Chloroethyl)ether	132-64-9	Dibenzofuran
95-57 <b>-</b> 8	2-Chlorophenol	121-14-2	2,4-Dinitrotoluene
541-73-1	1,3-Dichlorobenzene	84-66-2	Diethylphthalate
106-46-7	1,4-Dichlorobenzene	7005-72-3	4-Chlorophenyl-phenylether
95-50-1	1,2-Dichlorobenzene	86-73-7	Fluorene
95-48-7	2-Methylphenol	100-01-6	4-Nitroaniline
108-60-1	bis(2-Chloroisopropyl)ether	534-52-1	4,6-Dinitro-2-methylphenol
106-44-5	3 & 4-Methylphenol	86-30-6	N-Nitrosodiphenylamine
621-64-7	N-Nitroso-di-n-propylamine	126-73-8	Tri-n-butyl phosphate
67-72-1	Hexachloroethane	103-33-3	Azobenzene
98-95-3	Nitrobenzene	101-55-3	4-Bromophenyl-phenylether
78-59-1	Isophorone	118-74-1	Hexachlorobenzene
88-75-5	2-Nitrophenol	87-86-5	Pentachlorophenol
105-67-9	2,4-Dimethylphenol	85-01-8	Phenanthrene
111-91-1	bis(2-Chloroethoxy)methane)	120-12-7	Anthracene
120-83-2	2,4-Dichlorophenol	86-74-8	Carbazole
120-82-1	1,2,4-Trichlorobenzene	84-74-2	Di-n-butylphthalate
91-20-3	Naphthalene	206-44-0	Fluoranthene
106-47-8	4-Chloroaniline	129-00-0	Pyrene
87-68-3	Hexachlorobutadiene	85-68-7	Butylbenzylphthalate
59-50-7	4-Chloro-3-methylphenol	91-94-1	3,3'-Dichlorobenzidine
91-57-6	2-Methylnaphthalene	218-01-9	Chrysene
77-47-4	Hexachlorocyclopentadiene	56-55-3	Benzo(a)anthracene
88-06-2	2,4,6-Trichlorophenol	117-81-7	bis(2-Ethylhexyl)phthalate
95-95-4	2,4,5-Trichlorophenol	117-84-0	Di-n-octylphthalate
91-58-7	2-Chloronaphthalene	205-99-2	Benzo(b)fluoranthene
88-74-4	2-Nitroaniline	207-08-9	Benzo(k)fluoranthene
131-11-3	Dimethylphthalate	50-32-8	Benzo(a)pyrene
606-20-2	2,6-Dinitrotoluene	193-39-5	Indeno(1,2,3-cd)pyrene
208-96-8	Acenaphthylene	53-70-3	Dibenzo(a,h)anthracene
99-09-2	3-Nitroaniline	191-24-2	Benzo(g,h,I)perylene

FEED SAMPLES ANALYSIS REPORTS

FINAL REPORT for 150WM:181 198

Log Type: \*\* RCRA - PLANT \*\*

Log Number : 00-09272 Phone Number : 6-3226 Report for : NWCFMailstop : 5116

Date Approved : Feb 21 2001 Time Approved : 09:01 Date Received : Sep 27 2000

Time Received : 10:38

GWA charged : 561211110 Reviewed by BRIAN STORMS

MSA mR/hr : ? Signature

Hazard Index : >1E4 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_

COMMENTS:

Lab Field

Analysis Spl ID Spl ID Method Analyst Results \_\_\_\_\_

Total Sr 0CD15 150WM/NEOPRENE 23381 BJS 7.549E+05 +- 4.0E+03 D/S/ML Tritium 0CD15 150WM/NEOPRENE 33011 WDT 5.85E+02 +- 5.0E+01 D/S/ML End of Report -- 2 results.

## F I N A L R E P O R T for 150FEED:103 198/181

Log Type: \*\* PLANT \*\*

Log Number : 00-09274
Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Feb 20 2001 Time Approved : 10:11 Date Received : Sep 27 2000

Time Received : 11:11

Reviewed by TIFFANY PARK GWA charged : 561211110

Signature \_\_\_\_\_ MSA mR/hr : COLD

Hazard Index : <1E4 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

COMMENTS:

	Lab	Field	
Analysis	Spl ID	Spl ID	Method Analyst Results
			FROM DRU 1 FC4F-00 - 0 OF 02 Normal Agid
Acid	0CD21	150WM/NEOPRENE	57012 RAH 1.564E+00 +- 8.8E-02 Normal Acid
Aluminum	0CD25	FEED150/PLASTIC	87100 BCS 2.209E-01 +- 3.2E-03 MOLAR
Boron	0CD25	FEED150/PLASTIC	87100 BCS 1.30E-02 +- 2.9E-03 MOLAR
Cadmium	0CD25	FEED150/PLASTIC	87100 BCS 4.11E-03 +- 8.8E-04 MOLAR
Calcium	0CD25	FEED150/PLASTIC	87100 BCS 4.376E-02 +- 7.4E-04 MOLAR
Chloride	0CD21	150WM/NEOPRENE	57171 AWO 6.34E+02 +- 2.0E+01 ug/mL
Chromium	0CD25	FEED150/PLASTIC	87100 BCS 2.75E-03 +- 7.0E-04 MOLAR
Co60	0CD21	150WM/NEOPRENE	83993 SJH 2.61E+03 +- 1.3E+02 d/s/ml
Cs134	0CD21	150WM/NEOPRENE	83993 SJH 2.61E+03 +- 1.4E+02 d/s/ml
Cs137	0CD21	150WM/NEOPRENE	83993 SJH 9.56E+05 +- 3.5E+04 d/s/ml
Eu154	0CD21	150WM/NEOPRENE	83993 SJH 7.91E+03 +- 5.0E+02 d/s/ml
Eu155	0CD21	150WM/NEOPRENE	83993 SJH 1.79E+03 +- 2.2E+02 d/s/ml
Fluoride	0CD21	150WM/NEOPRENE	57093 BCS 1.612E+03 +- 1.8E+01 ug/mL
Iron	0CD25	FEED150/PLASTIC	87100 BCS 1.13E-02 +- 1.4E-03 MOLAR
Mercury	0CD25	FEED150/PLASTIC	87802 RDW 3.98E+02 +- 2.5E+01 ug/ml
NB94	0CD21	150WM/NEOPRENE	83993 SJH 1.17E+02 +- 1.2E+01 d/s/ml
Nitrate	0CD21	150WM/NEOPRENE	97074 BCS 2.839E+00 +- 2.7E-02 Molar
Potassium	0CD25	FEED150/PLASTIC	12800 SDN 3.21 E+04 ug/mL
Sodium	0CD25	FEED150/PLASTIC	12800 SDN 1.31 E+05 ug/mL
SpGr	0CD21	150WM/NEOPRENE	47981 AWO 1.16019E+00 +- 2.7E-04 @ 25/4
Sulfate	0CD21	150WM/NEOPRENE	97168 BCS 3.04E+03 +- 8.4E+02 ug/ml
Uranium PreP	0CD21	150WM/NEOPRENE	17929 BGP 1.0E+00 ml
Zirconium	0CD25	FEED150/PLASTIC	87100 BCS 5.3E-03 +- 2.9E-03 MOLAR
Zr95	0CD21	150WM/NEOPRENE	83993 SJH $4.63E+02 +- 3.4E+01 d/s/ml$
End of Report	24 r	esults.	

#### FINAL REPORT for 150FEED:103 199

Log Type: \*\* PLANT \*\*

Log Number : 00-10022 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Mar 28 2001 Time Approved : 10:26 Date Received : Oct 12 2000

Time Received : 15:33

Reviewed by KIMBERLY HONAS GWA charged : 561211290

Signature \_\_\_\_\_ MSA mR/hr : CELL

Hazard Index : 1E8 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

COMMENTS:

	Lab	Field	
Analysis	Spl ID	Spl ID	Method Analyst Results
Acid	0CD94	FEED150/NEOPRENE	57012 AWO 1.596E+00 +- 8.8E-02 Normal Acid
Aluminum		FEED150/PLASTIC	87100 NWJ 2.189E-01 +- 6.3E-03 MOLAR
	0CD95	FEED150/PLASTIC	87100 NWJ 1.34E-02 +- 2.9E-03 MOLAR
Boron Cadmium	0CD95	FEED150/PLASTIC	87100 NWJ 4.26E-03 +- 8.8E-04 MOLAR
Calcium	0CD95	FEED150/PLASTIC	87100 NWJ 4.526E-02 +- 7.4E-04 MOLAR
	0CD95 0CD94	FEED150/PLASTIC FEED150/NEOPRENE	57171 AWO 4.15E+02 +- 1.7E+01 ug/mL
Chloride	0CD94 0CD95	FEED150/PLASTIC	87100 NWJ 2.52E-03 +- 7.0E-04 MOLAR
Chromium	0CD95 0CD94	FEED150/PLASTIC FEED150/NEOPRENE	93993 SJH 2.40E+03 +- 2.0E+02 d/s/ml
C060		•	93993 SJH 2.35E+03 +- 1.4E+02 d/s/ml
Cs134	0CD94	FEED150/NEOPRENE	
Cs137	0CD94	FEED150/NEOPRENE	
Eul54	0CD94	FEED150/NEOPRENE	93993 SJH 8.28E+03 +- 8.1E+02 d/s/ml
Eu155	0CD94	FEED150/NEOPRENE	93993 SJH 1.67E+03 +- 2.4E+02 d/s/ml
Fluoride	0CD94	FEED150/NEOPRENE	57093 BGP 2.594E+03 +- 3.8E+01 ug/mL
Iron	0CD95	FEED150/PLASTIC	87100 NWJ 1.21E-02 +- 1.4E-03 MOLAR
Mercury	0CD95	FEED150/PLASTIC	87802 BET 2.05E+02 +- 2.6E+01 ug/ml
NB94	0CD94	FEED150/NEOPRENE	93993 SJH 1.140E+02 +- 9.9E+00 d/s/ml
Nitrate	0CD94	FEED150/NEOPRENE	97074 BET 3.111E+00 +- 2.7E-02 Molar
Potassium	0CD95	FEED150/PLASTIC	12800 RHH 4.86E+03 ug/mL
Sb125	0CD94	FEED150/NEOPRENE	93993 SJH 1.85E+03 +- 1.8E+02 d/s/ml
Sodium	0CD95	FEED150/PLASTIC	12800 RHH 1.95E+04 ug/mL
SpGr	0CD94	FEED150/NEOPRENE	47981 AWO 1.15490E+00 +- 2.7E-04 @ 25/4
Sulfate	0CD94	FEED150/NEOPRENE	97168 BET 3.43E-02 +- 8.7E-03 molar
Uranium	0CD94	FEED150/NEOPRENE	17920 RAH 7.5E-02 +- 1.4E-02 G/L
Zirconium	0CD95	FEED150/PLASTIC	87100 NWJ 5.4E-03 +- 2.9E-03 MOLAR
Zr95		FEED150/NEOPRENE	93993 SJH 4.27E+02 +- 5.2E+01 d/s/ml
End of Report	25 r		

#### FINAL REPORT for 150WM:181 199

Log Type: \*\* RCRA - PLANT \*\*

Log Number : 00-100415 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Feb 21 2001 Time Approved : 09:00 Date Received : Oct 05 2000

Time Received : 11:35

Reviewed by BRIAN STORMS GWA charged : 591211290

MSA mR/hr : HOT Signature \_\_\_\_

Hazard Index : >1E4 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS:

Lab Field Analysis Spl ID Spl ID

Method Analyst Results

Total Sr 0CF26 150WM/NEOPRENE 23381 BJS 8.983E+05 +- 4.3E+03 D/S/ML
Tritium 0CF26 150WM/NEOPRENE 33011 WDT 5.71E+02 +- 4.9E+01 D/S/ML
End of Report -- 2 results.

FINAL REPORT for 150WM:181 200

Log Type: \*\* RCRA - PLANT \*\*

Log Number : 00-10164 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Feb 21 2001 Time Approved : 08:59 Date Received : Oct 17 2000

Time Received: 17:04

Reviewed by BRIAN STORMS GWA charged : 561211290

MSA mR/hr : CELL Signature \_\_\_\_

Hazard Index : 1E8 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_

COMMENTS:

Analysis Spl ID Spl ID Method Analyst Results

Total Sr 0CG62 WM181/NEOPREN 23381 BJS 7.656E+05 +- 4.3E+03 D/S/ML

Tritium 0CG62 WM181/NEOPREN 33011 WDT 4.66E+02 +- 4.0E+01 D/S/ML

End of Report -- 2 results.

#### FINAL REPORT for 150FEED:103 200

Log Type: \*\* PLANT \*\*

Report for : NWCF Mailstop : 5116 Log Number : 00-10167 Phone Number : 6-3226

Date Approved : Nov 02 2000 Time Approved : 10:54 Date Received : Oct 17 2000

Time Received: 05:53

GWA charged : 561211290 Reviewed by JEFF LAUG

MSA mR/hr : COLD Signature \_\_\_\_

Lab QC/QA reviewed by Hazard Index : <1E4

PCBs >50 ppm : NO Signature \_\_\_\_\_

	Lab	Field		
Analysis		Spl ID	Method An	alyst Results
Acid	0CG70	103	57012 RA	LH 1.633E+00 +- 8.8E-02 Normal Acid
Aluminum	0CG70	103	87100 BC	
Aluminum	00070	0CG74	87100 BC	
Boron	0CG70	103	87100 NW	
Cadmium	0CG70	103	87100 NW	
		103	87100 NW	
Calcium	0CG70		57171 RA	
Chloride	0CG70	103	87100 NW	——————————————————————————————————————
Chromium	0CG70	103		
Co60	0CG70	103	93993 SJ	, ,
Cs134	0CG70	103	93993 SJ	• •
Cs137	0CG70	103	93993 SJ	
Eu154	0CG70	103	93993 SJ	
Eu155	0CG70	103	93993 SJ	
Fluoride	0CG70	103		S 1.543E+03 +- 8.4E+01 ug/mL
Iron	0CG70	103	87100 NW	
Mercury		0CG74		H 1.21E+02 +- 2.7E+01 ug/ml
NB94	0CG70	103	93993 SJ	
Nitrate	0CG70	103	97074 BG	P 3.222E+00 +- 2.7E-02 Molar
Potassium		0CG74	12800 RH	[H 4.97E+03 ug/mL
Sb125	0CG70	103	93993 SJ	H 1.36E+03 +- 1.2E+02 d/s/ml
Sodium		0CG74	12800 RH	[H 1.96E+04 ug/mL
SpGr	0CG70	103	47981 RA	H 1.15786E+00 +- 2.7E-04 @ 25/4
Sulfate	0CG70	103	97168 BG	P 3.70E+03 +- 8.5E+02 ug/ml
Uranium	0CG70	103	17920 RA	H 7.4E-02 +- 1.4E-02 G/L
Zirconium	0CG70	103	87100 NW	J 5.3E-03 +- 2.8E-03 MOLAR
Zr95	0CG70	103	93993 SJ	H 4.49E+02 +- 5.2E+01 d/s/ml
End of Report	26 r	esults.		

#### FINAL REPORT for 150FEED:184

Log Type: \*\* PLANT \*\*

Log Number : 01-04103 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Apr 24 2001 Time Approved : 15:44 Date Received : Apr 10 2001

Time Received: 13:42

GWA charged : 561211290 Reviewed by KIMBERLY HONAS

MSA mR/hr : HOT Signature \_\_\_\_

Hazard Index : <1E4 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

	Lab	Field		
Analysis	Spl ID	Spl ID	Method Analyst Results	_
Acid	1AU03	FEED150/NEOPRENE	57012 RAH 1.842E+00 +- 5.5E-02 Normal Acid	
Aluminum	1AU03	FEED150/NEOPRENE	87100 RAH 5.26E-01 +- 2.4E-02 MOLAR	
Boron		FEED150/PLASTIC	87100 RAH 7.6E-03 +- 1.4E-03 MOLAR	
Cadmium		FEED150/PLASTIC	87100 RAH < 9.47983E-04 MOLAR	
Calcium		FEED150/PLASTIC	87100 RAH 1.618E-02 +- 3.8E-04 MOLAR	
Chloride	1AU03	FEED150/NEOPRENE	57171 RAH 9.77E+02 +- 1.8E+01 ug/mL	
Chromium	1AU04	FEED150/PLASTIC	87100 RAH 1.69E-03 +- 4.7E-04 MOLAR	
Co57	1AU03	FEED150/NEOPRENE	43993 MLE 1.171E+04 +- 7.2E+02 pC/ml	
Co60	1AU03	FEED150/NEOPRENE	43993 MLE 1.357E+04 +- 6.0E+02 pC/ml	
Cs137	1AU03	FEED150/NEOPRENE	43993 MLE 1.280E+07 +- 2.8E+05 pC/ml	
Eu154	1.AU03	FEED150/NEOPRENE	43993 MLE 2.99E+04 +- 3.5E+03 pC/ml	
Fluoride	1AU03	FEED150/NEOPRENE	57093 BGP 5.52E+02 +- 4.3E+01 ug/mL	
Iron	1AU04	FEED150/PLASTIC	87100 RAH 1.422E-02 +- 8.5E-04 MOLAR	
Mercury	1AU04	FEED150/PLASTIC	87802 RDW 1.50E+02 +- 1.2E+01 ug/ml	
Nitrate	1AU03	FEED150/NEOPRENE	97074 BCS 3.91E+00 +- 1.2E-01 Molar	
PREP	1AU03	FEED150/NEOPRENE	17961 BCS Prep Completed 2001-04-12 10:07	
Potassium	1AU04	FEED150/PLASTIC	12800 SDN 3.50 E+03 ug/mL	
Sodium	1AU04	FEED150/PLASTIC	12800 SDN 3.04 E+04 ug/mL	
SpGr	1AU03	FEED150/NEOPRENE	47981 RAH 1.22196E+00 +- 3.7E-04 @ 25/4	
Sulfate	1AU03	FEED150/NEOPRENE	97168 BCS	
Uranium	1AU03	FEED150/NEOPRENE	17920 RAH 4.41E-02 +- 1.7E-03 G/L	
Zirconium	1AU04	FEED150/PLASTIC	87100 RAH < 2.78182E-03 MOLAR	
End of Report	22 re	sults.		

#### FINAL REPORT for 150FEED:184 281

Log Type: \*\* PLANT \*\*

Log Number : 01-04125 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 01 2001 Time Approved : 08:15 Date Received : Apr 13 2001

Time Received: 00:15

Reviewed by TIFFANY PARK GWA charged : 561211290

Signature \_\_\_\_ MSA mR/hr : COLD

Hazard Index : <1E4 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_\_

Analysis		Field Spl ID	Method Analyst Results
Acid	1AU88	FEED150/NEOPRENE	57012 RDW 1.823E+00 +- 1.9E-02 Normal Acid
Aluminum	1AU89	FEED150/PLASTIC	87100 BCS 5.37E-01 +- 2.0E-02 MOLAR
	1AU89	FEED150/PLASTIC	87100 BCS > 1.34297E-01 MOLAR
Boron	1AU89	FEED150/PLASTIC	87100 BCS 8.2E+01 +- 1.5E+01 UG/ML
Cadmium	1AU89	FEED150/PLASTIC	87100 BCS < 1.06553E+02 UG/ML
Calcium	1AU89	FEED150/PLASTIC	87100 BCS 6.53E+02 +- 1.5E+01 UG/ML
Chloride	1AU88	FEED150/NEOPRENE	57171 BCS 9.87E+02 +- 1.8E+01 ug/mL
Chromium	1AU89	FEED150/PLASTIC	87100 BCS 8.6E+01 +- 2.4E+01 UG/ML
Co60	1AU88	FEED150/NEOPRENE	
Cs134	1AU88	FEED150/NEOPRENE	83993 SJH 5.46E+06 +- 3.2E+05 pC/l
Cs137	1AU88	FEED150/NEOPRENE	83993 SJH 1.424E+10 +- 4.6E+08 pC/l
Eul54	1AU88	FEED150/NEOPRENE	83993 SJH 3.19E+07 +- 2.5E+06 pC/l
Fluoride	1AU88	FEED150/NEOPRENE	57093 AWO 4.90E+02 +- 4.1E+01 ug/mL
Iron	1AU89	FEED150/PLASTIC	87100 BCS 7.72E+02 +- 4.8E+01 UG/ML
Mercury	1AU89	FEED150/PLASTIC	87802 RDW 1.27E+02 +- 1.2E+01 ug/ml
Nitrate	1AU88	FEED150/NEOPRENE	97074 BCS 3.89E+00 +- 1.2E-01 Molar
Potassium	1AU89	FEED150/PLASTIC	12800 RHH 3.44E+03 ug/mL
Sodium	1AU89	FEED150/PLASTIC	12800 RHH 3.02E+04 ug/mL
SpGr	1AU88	FEED150/NEOPRENE	47981 BCS 1.22008E+00 +- 3.7E-04 @ 25/4
Sulfate	1AU88	FEED150/NEOPRENE	97168 BCS 8.4E+02 +- 1.1E+02 ug/ml
Uranium	1AU88	FEED150/NEOPRENE	17920 BCS 4.40E-02 +- 1.7E-03 G/L
Zirconium	1AU89	FEED150/PLASTIC	87100 BCS < 2.53758E+02 UG/ML
	22 r		

#### FINAL REPORT for 150FEED:184 WM184

Log Type: \*\* PLANT \*\*

Report for : NWCF Mailstop : 5116 Log Number : 01-04142 Phone Number : 6-3226

Date Approved : May 01 2001 Time Approved : 08:16 Date Received : Apr 14 2001 Time Received : 12:25

Reviewed by TIFFANY PARK GWA charged : 561211290

MSA mR/hr : CELL Signature \_\_\_\_\_

Lab QC/QA reviewed by Hazard Index : 1E7

Signature \_\_\_\_\_ PCBs >50 ppm : NO

	Lab	Field	
Analysis	Spl II	Spl ID	Method Analyst Results
Acid	1AU93	FEED150/NEOPRENE	57012 RDW 1.854E+00 +- 1.9E-02 Normal Acid
Aluminum	1AU94	FEED150/PLASTIC	87100 BCS 5.21E-01 +- 2.0E-02 MOLAR
Am241	1AU93	FEED150/NEOPRENE	83993 SJH 1.30E+08 +- 1.4E+07 pC/l
Boron	1AU93	FEED150/NEOPRENE	87100 BCS 8.9E+01 +- 1.5E+01 UG/ML
Cadmium	1AU93	FEED150/NEOPRENE	87100 BCS < 1.06553E+02 UG/ML
Calcium	1AU93	FEED150/NEOPRENE	87100 BCS 6.77E+02 +- 1.5E+01 UG/ML
Chloride	1AU93	FEED150/NEOPRENE	57171 BCS 1.005E+03 +- 1.8E+01 ug/mL
Chromium	1AU93	FEED150/NEOPRENE	87100 BCS 9.2E+01 +- 2.4E+01 UG/ML
Co60	1AU93	FEED150/NEOPRENE	83993 SJH 1.697E+07 +- 7.9E+05 pC/l
Cs134	1AU93	FEED150/NEOPRENE	83993 SJH 5.55E+06 +- 3.1E+05 pC/l
Cs137	1AU93	FEED150/NEOPRENE	83993 SJH 1.438E+10 +- 5.5E+08 pC/l
Eu154	1AU93	FEED150/NEOPRENE	83993 SJH 3.24E+07 +- 2.8E+06 pC/l
Fluoride	1AU93	FEED150/NEOPRENE	57093 AWO 4.94E+02 +- 4.2E+01 ug/mL
Iron	1AU93	FEED150/NEOPRENE	87100 BCS 8.32E+02 +- 4.8E+01 UG/ML
Mercury	1AU94	FEED150/PLASTIC	87802 RAH 1.36E+02 +- 1.2E+01 ug/ml
Nitrate	1AU93	FEED150/NEOPRENE	97074 BCS 3.96E+00 +- 1.2E-01 Molar
Potassium	1AU94	FEED150/PLASTIC	12800 RHH 3.44E+03 ug/mL
Sodium	1AU94	FEED150/PLASTIC	12800 RHH 3.12E+04 ug/mL
SpGr	1AU93	FEED150/NEOPRENE	47981 AWO 1.24092E+00 +- 4.0E-04 @ 25/4
Sulfate	1AU93	FEED150/NEOPRENE	97168 BCS 1.02E+03 +- 1.1E+02 ug/ml
Uranium	1AU93	FEED150/NEOPRENE	17920 RAH 3.99E-02 +- 1.5E-03 G/L
Zirconium		FEED150/NEOPRENE	87100 BCS < 2.53758E+02 UG/ML
		esults.	
Tild Of Report	22 1		

#### FINAL REPORT for 150FEED:101 294

Log Type: \*\* PLANT \*\*

Log Number : 01-05062 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 24 2001 Time Approved : 13:26 Date Received : May 06 2001

Time Received: 15:45

Reviewed by TIFFANY PARK GWA charged : 561211290

MSA mR/hr : CELL Signature \_\_\_\_

Hazard Index : 1E8 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_

7 1		Field	Method Analyst Results
Analysis	Spr in	Spl ID	method Analyst Results
Acid	1BB06	FEED150/NEOPRENE	57012 RNR 1.781E+00 +- 5.4E-02 Normal Acid
Aluminum	1BB07	FEED150/PLASTIC	87100 BCS 4.51E-01 +- 2.4E-02 MOLAR
Boron	1BB07	FEED150/PLASTIC	87100 BCS 1.05E+02 +- 1.5E+01 UG/ML
Cadmium	1BB07	FEED150/PLASTIC	87100 BCS 1.75E+02 +- 4.8E+01 UG/ML
Calcium	1BB07	FEED150/PLASTIC	87100 BCS 1.013E+03 +- 1.5E+01 UG/ML
Chloride	1BB06	FEED150/NEOPRENE	57171 RNR 8.02E+02 +- 1.7E+01 ug/mL
Chromium	1BB07	FEED150/PLASTIC	87100 BCS 1.01E+02 +- 2.4E+01 UG/ML
Cs134	1BB06	FEED150/NEOPRENE	93993 SJH 2.12E+07 +- 2.1E+06 pC/l
Cs137	1BB06	FEED150/NEOPRENE	93993 SJH 1.94E+10 +- 1.3E+09 pC/l
Fluoride	1BB06	FEED150/NEOPRENE	57093 BCS 9.49E+02 +- 6.0E+01 ug/mL
Iron	1BB07	FEED150/PLASTIC	87100 BCS 7.42E+02 +- 4.8E+01 UG/ML
Mercury	1BB07	FEED150/PLASTIC	87802 RAH 1.23E+02 +- 2.9E+01 ug/ml
NB94	1BB06	FEED150/NEOPRENE	93993 SJH 2.09E+06 +- 2.8E+05 pC/l
Nitrate	1BB06	FEED150/NEOPRENE	97074 BGP 3.279E+00 +- 4.5E-02 Molar
PREP	1BB06	FEED150/NEOPRENE	17961 BCS Prep Completed 2001-05-15 08:46
Potassium	1BB07	FEED150/PLASTIC	12800 RHH 3.86E+03 ug/mL
Sodium	1BB07	FEED150/PLASTIC	12800 RHH 2.69E+04 ug/mL
SpGr	1BB06	FEED150/NEOPRENE	47981 BGP 1.19761E+00 +- 3.4E-04 @ 25/4
Sulfate	1BB06	FEED150/NEOPRENE	97168 BGP 2.77E+03 +- 4.0E+02 ug/ml
Uranium	1BB06	FEED150/NEOPRENE	17920 BCS 4.90E-02 +- 2.9E-03 G/L
Zirconium	1BB07	FEED150/PLASTIC	87100 BCS < 2.54584E+02 UG/ML
Zr95	1BB06	FEED150/NEOPRENE	93993 SJH 4.77E+06 +- 7.4E+05 pC/l
End of Report	22 r	esults.	

FINAL REPORT for 150WM:101 294

Log Type: \*\* RCRA - PLANT \*\*

Log Number : 01-05063 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 30 2001 Time Approved : 13:13 Date Received : May 06 2001

Time Received: 15:55

GWA charged : 561211290 Reviewed by TIFFANY PARK

MSA mR/hr : CELL Signature \_\_\_\_\_

Hazard Index : 1E8 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS:

Analysis Spl ID Spl ID Method Analyst Results

PREP 1BB08 150WM/NEOPRENE 17961 BGP Prep Completed 2001-05-16 06:07

Total Sr 1BB08 150WM/NEOPRENE 23381 BJS 2.50E+05 +- 3.6E+04 D/S/ML

Tritium 1BB08 150WM/NEOPRENE 33011 WDT 8.26E+02 +- 7.1E+01 D/S/ML

End of Report -- 3 results.

FINAL REPORT for 150WM:101 298

Log Type: \*\* RCRA - PLANT \*\*

Log Number : 01-05112 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 30 2001 Time Approved : 13:13 Date Received : May 11 2001

Time Received : 09:43

Reviewed by TIFFANY PARK GWA charged : 561211290

MSA mR/hr : ? Signature \_\_\_\_

Lab QC/QA reviewed by Hazard Index : >1E4

Signature \_\_\_\_\_ PCBs >50 ppm : NO

COMMENTS:

Lab Field Analysis Spl ID Spl ID Method Analyst Results

PREP 1BD35 150WM/NEOPRENE 17961 BCS Prep Completed 2001-05-15 08:48
Total Sr 1BD35 150WM/NEOPRENE 23381 BJS 6.19E+05 +- 6.5E+04 D/S/ML
Tritium 1BD35 150WM/NEOPRENE 33011 WDT 8.11E+02 +- 6.9E+01 D/S/ML
End of Report -- 3 results.

#### INTERIM REPORT for BOPR: NCC-101

Log Type: \*\* RCRA \*\*

Log Number : 01-06071 Phone Number : 6-7552 Report for : JD LONG Mailstop : 5218

Date Received : Jun 07 2001 Time Received : 09:00

GWA charged : 561C022A9

MSA mR/hr : HOT

Hazard Index : >1E4

PCBs >50 ppm : NO

COMMENTS: If fail to get a acid result in range then run pH. pH does not

need to be RCRA

		Spl ID	Method Analyst Results
Agid			57012 RNR 1.711E+00 +- 5.4E-02 Normal Acid
Aluminum	1B126	BP10120101	42900 LAM 1.14534E+07 ug/L
Antimony	1BT26	BP10120101	42900 LAM Not Detected: IDL= 235 ug/L
Arsenic	1BT26	BP10120101	42900 LAM Not Detected: IDL= 235 ug/L 42900 LAM 3.35E+02 ug/L
Barium			42900 LAM 3.985E+03 ug/L
Beryllium	1BT26	BP10120101	
Cadmium		BP10120101	
Chromium		BP10120101	
Cobalt	1BI26	BP10120101	
Copper		BP10120101	
Fluoride	1BI26	BP10120101	57093 AWO 8.26654E+02 ug/mL
Lead	1BI26	BP10120101	42900 LAM 1.1125E+05 ug/L
Manganese	1BI26	BP10120101	42900 LAM 4.30765E+05 ug/L
Mercury	1BI26	BP10120101	12800 SDN 1.49E+05 ug/L
Nickel	1BI26	BP10120101	42900 LAM 6.64E+04 ug/L
SVOA (TOTAL)	1BI26	BP10120101	9270
Selenium	1BI26	BP10120101	42900 LAM Not Detected: IDL= 240 ug/L
Silver	1BI26	BP10120101	42900 LAM 1.2E+02 ug/L
TIC	1BI26	BP10120101	18060 RDW MDL=119.004 ug/ml
TOC		BP10120101	18060 RDW 6.08462E+02 ug/ml
Thallium			42900 LAM Not Detected: IDL= 200 ug/L
UDS	1BI26	BP10120101	17972 BCS 0.619 g(UDS)/ L
Uranium		BP10120101	17920 BCS 4.98E-02 +- 2.9E-03 G/L
VOA (TOTAL)	1BI26	BP10120101	9260
		BP10120201	
Vanadium	1BI26	BP10120101	42900 LAM 6.1E+02 ug/L
Zinc	1BI26	BP10120101	42900 LAM 6.1E+02 ug/L 42900 LAM 4.12784E+04 ug/L
End of Report	27 re	esults.	

CONCENTRATED BOTTOMS SAMPLES ANALYSIS REPORTS

#### F I N A L R E P O R T for 150B0T119 296/297

Log Type: \*\* PLANT \*\*

Log Number : 01-05106 Phone Number : 6-3226 Report for : NWCF Mailstop : 5216

Date Approved : Feb 04 2002 Time Approved : 13:07 Date Received : May 10 2001 Time Received : 23:55

GWA charged : 561211295 Reviewed by BRIAN STORMS

MSA mR/hr : HOT Signature \_\_\_\_

Hazard Index : 1E8 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

	Lab	Field			
Analysis	Spl ID	Spl ID	Method	Ana	lyst Results
Acid	1BD27	150BOT119PLAS	57012	BGP	2.559E+00 +- 6.0E-02 Normal Acid
Aluminum		150BOT119PLAS			5.18E-01 +- 2.4E-02 MOLAR
	1BD27	150BOT119PLAS	57171	BGP	8.02E+02 +- 1.7E+01 ug/mL
Co60	1BD28	150BOT119	93993	SJH	3.32E+07 +- 2.6E+06 pC/l
	1BE80	150BOT119 DUP	83993	SJH	3.71E+07 +- 1.8E+06 pC/l
Cs134	1BD28	150BOT119	93993	SJH	2.33E+07 +- 1.8E+06 pC/l
	1BE80	150BOT119 DUP	83993	SJH	2.38E+07 +- 1.2E+06 pC/l
Cs137	1BD28	150BOT119	93993	SJH	2.13E+10 +- 1.6E+09 pC/l
	1BE80	150BOT119 DUP			2.116E+10 +- 7.0E+08 pC/l
Eu154	1BD28	150BOT119	93993	SJH	8.42E+07 +- 7.8E+06 pC/l
	1BE80	150BOT119 DUP	83993	SJH	9.66E+07 +- 7.8E+06 pC/l
Fluoride	1BD27	150BOT119PLAS	57093	BGP	8.85E+02 +- 7.8E+01 ug/mL
Nitrate	1BD27	150BOT119PLAS	97074	BGP	4.03E+00 +- 1.2E-01 Molar
PREP	1BD27	150BOT119PLAS			Prep Completed 2001-05-11 02:41
	1BD28	150BOT119			Prep Completed 2001-05-15 08:47
Phosphorous	1BD27	150BOT119PLAS			1.19887E+02 ug/ml
Potassium	1BD28	150BOT119			6.20E+02 ug/mL
Sodium	1BD28	150BOT119			1.16E+04 ug/mL
SpGr	1BD27	150BOT119PLAS			1.07469E+00 +- 1.7E-04 @ 25/4
Tritium	1BD27	150BOT119PLAS			4.01E+02 +- 3.4E+01 D/S/ML
UDS	1BD27	150BOT119PLAS			5.3 G/L
Uranium	1BD28	150BOT119			5.87E-02 +- 2.4E-03 G/L
Zr95	1BD28	150BOT119			5.52E+06 +- 4.4E+05 pC/l
		150BOT119 DUP	83993	SJH	5.04E+06 +- 3.5E+05 pC/1
End of Report	24 r	esults.			

#### INTERIM REPORT for 150BOT119 324

Log Type: \*\* PLANT \*\*

Report for : NWCF Log Number : 01-06146 Mailstop : 5216 Phone Number : 6-3226

Date Received : Jun 15 2001

Time Received : 00:15

GWA charged : 561211295

MSA mR/hr : HOT

Hazard Index : 1E8

PCBs >50 ppm : NO

Analysis	Lab Field Spl ID Spl ID	Method Analyst Results
Acid Aluminum	1BK68 150BOT119 1BK68 150BOT119	57012 BGP 2.868E+00 +- 6.2E-02 Normal Acid 87100 BCS 8.59E-01 +- 3.0E-02 MOLAR
Chloride	1BK68 150BOT119	57171 BGP 1.016E+03 +- 1.8E+01 ug/mL
Fluoride I129	1BK68 150BOT119 1BK68 150BOT119	57093 AWO 1.71E+03 +- 1.2E+02 ug/mL 3539
Nitrate	1BK68 150BOT119 1BK68 150BOT119	97074 BCS 5.27E+00 +- 1.8E-01 Molar 17961 RAH Prep Completed 2001-07-02 13:24
PREP	1BK68 150BOT119 1BK68 150BOT119	17961 BCS Prep Completed 2001-06-17 17:06
Phosphorous Potassium	1BK68 150BOT119 1BK68 150BOT119	42900 RHH 4.33229E+02 ug/ml 12800 SDN 5.06E+03 ug/mL
Sodium	1BK68 150BOT119	12800 SDN 3.59E+04 ug/mL
SpGr Tritium	1BK68 150BOT119 1BK68 150BOT119	47981 BGP 1.35635E+00 +- 5.6E-04 @ 25/4 33011 WDT 5.6E+02 +- 1.0E+02 D/S/ML
UDS Uranium	1BK68 150BOT119 1BK68 150BOT119	17972 RNR 1.187 g/L 17920 RAH 7.39E-02 +- 3.7E-03 G/L
End of Report	15 results.	1/920 1011 /.392 02 1 3.72 03 072

#### INTERIM REPORT for BOPR: NCC-119

Log Type: \*\* RCRA \*\*

Report for : JD LONG Log Number : 01-06214 Mailstop : 5218 Phone Number : 6-7552

Date Received : Jun 21 2001

Time Received : 12:51

GWA charged : 561C022AA

MSA mR/hr : HOT

Hazard Index : >1E4

PCBs >50 ppm : NO

COMMENTS: If fail to get a acid result in range then run pH. pH does not

need to be RCRA samples to include trip blank (BP10130201)

	Lab			
Analysis	Spl ID	Spl ID	Method	Analyst Results
		BP10130101	57012	BET 1.741E+00 +- 5.4E-02 Normal Acid
Aluminum		BP10130101		
Antimony		BP10130101	42900	LAM 1.56E+03 ug/L
Arsenic		BP10130101	42900	LAM Not Detected: IDL= 580 ug/L
Barium		BP10130101		
Beryllium		BP10130101		
Cadmium		BP10130101		
Chromium	1BL27	BP10130101		
Cobalt	1BL27	BP10130101	42900	
Copper	1BL27	BP10130101	42900	
Fluoride	1BL27	BP10130101		
Lead	1BL27	BP10130101	42900	LAM 8.628E+04 ug/L
Manganese		BP10130101		
Mercury	1BL27	BP10130101	12800	SDN 1.14E+05 ug/L
Nickel		BP10130101	42900	LAM 5.532E+04 ug/L
SVOA (TOTAL)	1BL27	BP10130101	9270	
Selenium	1BL27	BP10130101	42900	LAM Not Detected: IDL= 960 ug/L
Silver	1BL27	BP10130101	42900	LAM Not Detected: IDL= 400 ug/L
TIC	1BL27	BP10130101	18060	RDW MDL=119.004 ug/ml
TOC	1BL27	BP10130101	18060	RDW 7.5499E+02 ug/ml
Thallium	1BL27			LAM Not Detected: IDL= 800 ug/L
UDS	1BL27	BP10130101	17972	RNR 1.288 g/L
Uranium	1BL27	BP10130101	17920	BCS 6.22E-02 +- 3.3E-03 G/L
VOA (TOTAL)	1BL27	BP10130101	9260	
	1BL28	BP10130201	9260	
Vanadium	1BL27	BP10130101	42900	LAM 5.6E+02 ug/L
Zinc		BP10130101		
End of Report				

#### INTERIM REPORT for 150BOT119 333/334

Log Type: \*\* PLANT \*\*

Report for : NWCF Log Number : 01-06233 Mailstop : 5216 Phone Number : 6-3226

Date Received : Jun 24 2001

Time Received : 11:42

GWA charged : 561211295

MSA mR/hr : CELL

Hazard Index : 1E8

PCBs >50 ppm : NO

Analysis	Lab Field Spl ID Spl ID	Method Analyst Results
Acid Aluminum Chloride Fluoride I129 Nitrate PREP	1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119	57012 RDW 2.830E+00 +- 6.2E-02 Normal Acid 87100 BCS 7.11E-01 +- 2.9E-02 MOLAR 57171 RDW 1.323E+03 +- 2.2E+01 ug/mL 57093 BGP 1.30E+03 +- 1.0E+02 ug/mL 3539 97074 BCS 7.22E+00 +- 1.7E-01 Molar 17961 RAH Prep Completed 2001-07-02 13:23
Phosphorous Potassium Sodium SpGr Tritium UDS Uranium End of Report	1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1BL41 150BOT119 1A results.	42900 RHH

**CONDENSED OVERHEADS SAMPLES ANALYSIS REPORTS** 

Log Type: \*\* PLANT \*\*

Log Number : 01-05061 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 06 2001 Time Approved : 17:51 Date Received : May 06 2001

Time Received: 12:17

Reviewed by W. (BILL) STRONG GWA charged : 561211296

MSA mR/hr : HOT Signature \_\_\_\_\_

Hazard Index : 1E5 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_

Analysis	Lab Spl ID	Field Spl ID	Method Analyst Results
AL/F RATIO	1BB04	150COND122/NEOPRENE	11023 BCS Ratio Not Performed
Acid	1BB04	150COND122/NEOPRENE	57012 RDW 4.67E-01 +- 1.2E-02 Normal Acid
Aluminum	1BB05	150COND122/PLASTIC	87100 BCS 1.08E-03 +- 2.3E-04 MOLAR
Chloride	1BB04	150COND122/NEOPRENE	57171 BCS 1.747E+02 +- 6.5E+00 ug/mL
Flash Point	1BB04	150COND122/NEOPRENE	17985 BCS NO FLASH @ 60.00 deg C corrected
Fluoride	1BB04	150COND122/NEOPRENE	57093 BCS Not Detected: MDL=7.757 ug/mL
GROSS BETA	1BB04	150COND122/NEOPRENE	87970 BCS 3.96E+05 +- 2.0E+04 B/Min/ml
Mercury	1BB05	150COND122/PLASTIC	87802 RDW 4.59E+00 +- 4.6E-01 ug/ml
Nitrate	1BB04	150COND122/NEOPRENE	97074 BCS 4.011E-01 +- 4.7E-03 Molar
SpGr	1BB04	150COND122/NEOPRENE	47981 BCS 1.01272E+00 +- 1.0E-04 @ 25/4
Sulfate	1BB04	150COND122/NEOPRENE	97168 BCS 1.47E+01 +- 3.9E+00 ug/ml
TOC	1BB04	150COND122/NEOPRENE	18060 RDW 1.517E+02 +- 9.1E+00 ug/ml
UDS	1BB04	150COND122/NEOPRENE	17972 BCS No Visible Solids.
Uranium	1BB04	150COND122/NEOPRENE	17920 BCS < 3.24219E-04 G/L
	1BB04	150COND122/NEOPRENE	17920 BCS < 3.24219E-04 G/L
End of Report	15 r	esults.	

# FINAL REPORT for 150COND122 294 Log Type: \*\* PLANT \*\*

Log Number : 01-05073 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 08 2001 Time Approved : 11:11 Date Received : May 07 2001 Time Received : 17:33

GWA charged : 561211296 Reviewed by CLAYNE GRIGG

MSA mR/hr : 1.0 Signature \_\_\_\_\_

Hazard Index : 1E5 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

	Lab	Field	
Analysis	Spl ID	Spl ID	Method Analyst Results
AL/F RATIO	1BB24	150COND122/NEOPRENE	11023 BGP Ratio Not Performed
Acid	1BB24	150COND122/NEOPRENE	57012 BGP 4.50E-01 +- 1.2E-02 Normal Acid
Aluminum	1BB25	150COND122/PLASTIC	87100 RAH < 7.55977E-04 MOLAR
Chloride	1BB24	150COND122/NEOPRENE	57171 BGP 1.72E+02 +- 1.1E+01 ug/mL
Flash Point	1BB24	150COND122/NEOPRENE	17985 BET NO FLASH @ 60.00 deg C corrected
Fluoride	1BB24	150COND122/NEOPRENE	57093 BGP Not Detected: MDL=6.982 ug/mL
GROSS BETA	1BB24	150COND122/NEOPRENE	87970 RAH 4.94E+04 +- 2.9E+03 B/Min/ml
Mercury	1BB25	150COND122/PLASTIC	87802 RAH 3.80E+00 +- 2.3E-01 ug/ml
Nitrate	1BB24	150COND122/NEOPRENE	97074 BGP 4.198E-01 +- 4.8E-03 Molar
SpGr	1BB24	150COND122/NEOPRENE	47981 BGP 1.01208E+00 +- 1.0E-04 @ 25/4
Sulfate	1BB24	150COND122/NEOPRENE	97168 BGP 1.15E+01 +- 4.0E+00 ug/ml
TOC	1BB24	150COND122/NEOPRENE	18060 BGP 1.313E+02 +- 9.1E+00 ug/ml
UDS	1BB24	150COND122/NEOPRENE	17972 BGP No Visible Solids.
Uranium	1BB24	150COND122/NEOPRENE	17920 RAH < 3.24219E-04 G/L
	1BB24	150COND122/NEOPRENE	17920 RAH < 3.24219E-04 G/L
End of Report	15 r	esults.	

Log Type: \*\* PLANT \*\*

Log Number : 01-05087
Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 09 2001 Time Approved : 02:13 Date Received : May 08 2001

Time Received : 14:53

Reviewed by BRIAN PASSMORE GWA charged : 561211296

MSA mR/hr : HOT Signature \_\_\_\_\_

Hazard Index : >1E4 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

Analysis	Lab Spl ID	Field Spl ID	Method Analyst Results	
AL/F RATIO Acid Aluminum	1BB54 1BB54 1BB55	150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/PLASTIC	11023 BCS Ratio Not Performed 57012 RNR 4.76E-01 +- 1.2E-02 Normal Acid 87100 BCS < 4.20091E-04 MOLAR	
Chloride	1BB54	150COND122/NEOPRENE	57171 RNR 1.828E+02 +- 6.6E+00 ug/mL	
Flash Point Fluoride	1BB54 1BB54 1BB54	150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE	17985 BET NO FLASH @ 60.00 deg C corrected 57093 BCS Not Detected: MDL=6.982 ug/mL 87970 RAH 5.01E+03 +- 7.3E+02 B/Min/ml	
GROSS BETA Mercury	1BB55	150COND122/PLASTIC	87802 RAH 3.73E+00 +- 2.3E-01 ug/ml	
Nitrate SpGr	1BB54 1BB54	150COND122/NEOPRENE 150COND122/NEOPRENE	97074 RAH 2.049E-02 +- 2.2E-04 Molar 47981 BET 1.01314E+00 +- 1.0E-04 @ 25/4	
Sulfate	1BB54	150COND122/NEOPRENE	97168 RAH < 2.86967E+00 ug/ml	
TOC	1BB54 1BB54	150COND122/NEOPRENE 150COND122/NEOPRENE	18060 BET 1.286E+02 +- 9.0E+00 ug/ml 17972 BET no visible solids	
Uranium	1BB54 1BB54	150COND122/NEOPRENE 150COND122/NEOPRENE	17920 BCS < 3.24219E-04 G/L 17920 BGP < 3.24219E-04 G/L	
End of Report		esults.	1,520 201 1 0.111111	

Log Type: \*\* PLANT \*\*

Log Number : 01-05092 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : May 10 2001 Time Approved : 01:06 Date Received : May 09 2001

Time Received: 18:47

GWA charged : 561211296 Reviewed by BRIAN PASSMORE

MSA mR/hr : HOT Signature \_\_\_\_\_

Hazard Index : >1E4 Lab QC/QA reviewed by

PCBs >50 ppm : NO Signature \_\_\_\_\_

	Lab	Field	Mathad	Analyst Dogulta
Analysis	SpI ID	Spl ID	меспоа	Analyst Results
AL/F RATIO	1BB93	150COND122/NEOPRENE		RAH Ratio Not Performed
Acid	1BB93	150COND122/NEOPRENE	57012	BGP 4.80E-01 +- 1.2E-02 Normal Acid
Aluminum	1BB94	150COND122/PLASTIC	87100	RAH < 7.51772E-04 MOLAR
Chloride	1BB93	150COND122/NEOPRENE	57171	BGP 1.86E+02 +- 1.1E+01 ug/mL
Flash Point	1BB93	150COND122/NEOPRENE	17985	BGP NO FLASH @ 60.00 deg C corrected
Fluoride	1BB93	150COND122/NEOPRENE	57093	BGP Not Detected: MDL=6.982 ug/mL
GROSS BETA	1BB93	150COND122/NEOPRENE	87970	RAH 3.52E+04 +- 2.2E+03 B/Min/ml
Mercury	1BB94	150COND122/PLASTIC	87802	RAH 2.31E+00 +- 2.3E-01 ug/ml
Nitrate	1BB93	150COND122/NEOPRENE	97074	BGP 4.233E-01 +- 4.7E-03 Molar
SpGr	1BB93	150COND122/NEOPRENE	47981	BGP 1.01307E+00 +- 1.0E-04 @ 25/4
Sulfate	1BB93	150COND122/NEOPRENE	97168	BGP 1.08E+01 +- 4.0E+00 ug/ml
TOC	1BB93	150COND122/NEOPRENE	18060	BGP 1.468E+02 +- 9.1E+00 ug/ml
UDS	1BB93	150COND122/NEOPRENE	17972	BGP No visible solids.
Uranium	1BB93	150COND122/NEOPRENE	17920	RAH < 3.24219E-04 G/L
	1BB93	150COND122/NEOPRENE	17920	RAH < 3.24219E-04 G/L
Deal of December	1	1 + .		

Log Type: \*\* PLANT \*\*

Log Number : 01-06145 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Jun 17 2001 Time Approved : 14:04 Date Received : Jun 14 2001

Time Received : 23:07

Reviewed by W. (BILL) STRONG GWA charged : 561211296

MSA mR/hr : HOT Signature \_\_\_\_\_

Hazard Index : >1E4 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

Analysis	Lab Spl ID	Field Spl ID	Method	Ana	lyst Results
AL/F RATIO	1BK66	150COND122/NEOPRENE			Ratio Not Performed
Acid	1BK66	150COND122/NEOPRENE	57012		
Aluminum	1BK67	150COND122/PLASTIC	•		< 4.20091E-04 MOLAR
Chloride	1BK66	150COND122/NEOPRENE	57171	BGP	1.986E+02 +- 6.8E+00 ug/mL
Flash Point	1BK66	150COND122/NEOPRENE			NO FLASH @ 60.00 deg C corrected
Fluoride	1BK66	150COND122/NEOPRENE	87092	BCS	Not Detected: MDL=3.504 ug/ml
GROSS BETA	1BK66	150COND122/NEOPRENE	87970	RAH	•
Mercury	1BK67	150COND122/PLASTIC	87802	RDW	5.49E+00 +- 2.4E-01 ug/ml
Nitrate	1BK66	150COND122/NEOPRENE	97074		
SpGr	1BK66	150COND122/NEOPRENE	47981	BGP	1.01377E+00 +- 1.0E-04 @ 25/4
Sulfate	1BK66	150COND122/NEOPRENE	97168	RAH	<b>3</b> .
TOC	1BK66	150COND122/NEOPRENE	18060	BGP	1.442E+02 +- 9.1E+00 ug/ml
UDS	1BK66	150COND122/NEOPRENE	17972	BCS	No Visible Solids
Uranium	1BK66	150COND122/NEOPRENE			< 3.24219E-04 G/L
	1BK66	150COND122/NEOPRENE	17920	RAH	< 3.24219E-04 G/L
End of Report	15 r	esults.			

#### INTERIM REPORT for BOPR:NCC-122

Log Type: \*\* RCRA \*\*

Report for : JD LONG Log Number : 01-06221 Mailstop : 5218 Phone Number : 6-7552

Date Received : Aug 13 2001

Time Received: 10:53

GWA charged : 561C022AB

MSA mR/hr : HOT

Hazard Index : >1E4

PCBs >50 ppm : NO

COMMENTS: If fail to get a acid result in range then run pH. pH does not

need to be RCRA

	Lab Field	
Analysis	spl ID spl ID	Method Analyst Results
Acid	1BL29 BP10140101	57012 BGP 5.14E-01 +- 1.2E-02 Normal Acid
Aluminum	1BL29 BP10140101	42900 LAM 1.703E+03 ug/L
Antimony	1BL29 BP10140101	42900 LAM Not Detected: IDL= 47 ug/L
Arsenic	1BL29 BP10140101	42900 LAM Not Detected: IDL= 29 ug/L
Barium	1BL29 BP10140101	42900 LAM 9.0E+00 ug/L
Beryllium	1BL29 BP10140101	42900 LAM Not Detected: IDL= 1 $ug/L$
Cadmium	1BL29 BP10140101	42900 LAM Not Detected: IDL= 4 ug/L
Chromium	1BL29 BP10140101	42900 LAM 2.7E+01 ug/L
Cobalt	1BL29 BP10140101	42900 LAM Not Detected: IDL= 10 ug/L
Copper	1BL29 BP10140101	42900 LAM Not Detected: IDL= 14 $ug/L$
Fluoride	1BL29 BP10140101	57093 BET 3.40346E+00 mg/L
Lead	1BL29 BP10140101	42900 LAM Not Detected: IDL= 63 ug/L
Manganese	1BL29 BP10140101	42900 LAM 1.3E+01 ug/L
Mercury	1BL29 BP10140101	12800 SDN 3950. ug/L
Nickel	1BL29 BP10140101	42900 LAM 3.2E+01 ug/L
SVOA (TOTAL)	1BL29 BP10140101	9270
Selenium	1BL29 BP10140101	42900 LAM Not Detected: IDL= 48 ug/L
Silver	1BL29 BP10140101	42900 LAM Not Detected: IDL= 20 ug/L
TIC	1BL29 BP10140101	18060 RDW mdl=23.8008 ug/ml
TOC	1BL29 BP10140101	18060 RDW 1.47853E+02 ug/ml
Thallium	1BL29 BP10140101	42900 LAM Not Detected: IDL= 40 ug/L
UDS	1BL29 BP10140101	17972 RNR 0.0 g/L
Uranium	1BL29 BP10140101	17920 BCS < 3.24219E-04 G/L
VOA (TOTAL)	1BL29 BP10140101	9260
	1BL30 BP10140201	9260
Vanadium	1BL29 BP10140101	42900 LAM Not Detected: IDL= 10 ug/L
Zinc	1BL29 BP10140101	42900 LAM 5.3E+01 ug/L
End of Report	27 results.	

Log Type: \*\* PLANT \*\*

Log Number : 01-06241 Phone Number : 6-3226 Report for : NWCF Mailstop : 5116

Date Approved : Jun 25 2001 Time Approved : 16:38 Date Received : Jun 24 2001

Time Received : 11:44

Reviewed by CLAYNE GRIGG GWA charged : 561211296

Signature \_\_\_\_ MSA mR/hr : CELL

Hazard Index : 1E5 Lab QC/QA reviewed by

Signature \_\_\_\_\_ PCBs >50 ppm : NO

Analysis	Lab Spl ID	Field Spl ID	Method Analyst Results
AL/F RATIO Acid Aluminum Chloride Flash Point Fluoride GROSS BETA Mercury Nitrate SpGr Sulfate TOC UDS Uranium	1BL43 1BL44 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43 1BL43	150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/PLASTIC 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/PLASTIC 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE 150COND122/NEOPRENE	11023 BCS Ratio Not Performed 57012 RDW 4.71E-01 +- 1.2E-02 Normal Acid 87100 BCS < 4.20091E-04 MOLAR 57171 RDW 1.91E+02 +- 1.1E+01 ug/mL 17985 BCS NO FLASH @ 60.00 deg C corrected 87092 BCS Not Detected: MDL=3.504 ug/ml 87970 BCS 6.90E+03 +- 7.9E+02 B/Min/ml 87802 RDW 3.27E+00 +- 4.6E-01 ug/ml 97074 BCS 4.007E-01 +- 9.0E-03 Molar 47981 BCS 1.01299E+00 +- 1.0E-04 @ 25/4 97168 BCS < 5.63559E+00 ug/ml 18060 RDW 1.888E+02 +- 9.3E+00 ug/ml 17972 BCS No Visible Solids 17920 BCS < 3.24219E-04 G/L 17920 BCS < 3.24219E-04 G/L
End of Report	1BL43 15 re	150COND122/NEOPRENE esults.	1/920 DC3 < 3.242196-04 G/H

#### LAB QUALIFIER LIST

For volatile and semi-volatile organic analytical results, the INEEL qualifiers to be used are as follows:

- U Indicates the compound was analyzed for but not detected. The sample quantification limit, or method detection limit (MDL) for EPA Method 524.2 (see Section C, Part I), must be corrected for dilution. For example, 10 U for phenol in water if the sample final volume is the protocol-specified final volume and the method quantification limit for phenol is  $10 \mu g/L$ . If a 1 to 10 dilution of extract is necessary, the reported limit is 100 U.
  - If an analyte is detected at a concentration that is less than the MDL, it shall be reported at the estimated quantification limit (EQL) (concentration of the low standard in the initial calibration) and a "U" flag shall be used on the Form I.
- Indicates an estimated value. This flag is used under the following circumstances: 1) either when estimating a concentration for tentatively identified compounds where a 1:1 response is assumed, 2) when the mass spectral data indicate the presence of a compound that meets the identification criteria but the result is less than the sample estimated quantification limit but greater than the method detection limit (MDL), and 3) when the retention time data indicates the presence of a compound that meets the pesticide/polychlorinated bisphenyls (PCBs) identification criteria and the result is less that the EQL but greater than zero. For example, if the sample quantification limit is 10 μg/L, but a concentration of 3 μg/L is calculated, report the result as 3 J. The sample estimated quantification limit must be adjusted for the U flag, so that if a sample with 24% moisture and a 1 to 10 dilution factor has a calculated concentration of 300 μg/kg and a sample quantification limit of 4300 μg/kg, report the concentration as 300 J on Form I. Note: The "J" code is not used and the compound is not reported as being identified for pesticide/PCBs results less than the Contract-Required Quantification Limit (CRQL), if the technical judgement of the pesticide residue analysis expert determines that the peaks used for compound identification resulted from instrument noise or other interferences (column bleed, solvent contamination, etc).
- N Indicates presumptive evidence of a compound. This flag is only used for tentatively identified compounds, where the identification is based on a mass spectral library search. The N flag is applicable only to tentatively identified compound results. For generic characterization of a tentatively identified compound, such as chlorinated hydrocarbon, the N code is not used.
- B This flag is used when the analyte is found in the associated blank as well as in the sample. It indicates possible/probable blank contamination and warns the data user to take appropriate action. This flag shall be used for a TIC as well as for a positively identified target compound. This flag shall not be used for flagging results on Forms I for the method blank analyses.
- E This flag identifies compounds whose concentrations exceed the calibration range of the gas chromatography (GC) or gas chromatography/mass spectrometry (GC/MS) instrument for that specific analysis. If one or more compounds has a response representing a concentration greater than the highest concentration used in the initial calibration of the instrument, the sample or extract shall be diluted and re-analyzed. All such compounds with a response greater than the highest concentration used in the initial calibration shall have the

concentration flagged with an "E" on the Form I for the original analysis. If the dilution of the extract causes any compounds identified in the first analysis to be below the calibration range in the second analysis, then the results of both analyses shall be reported on separate Forms I. The Form I for the diluted sample shall have the "DL" suffix appended to the sample number. NOTE: For total xylenes, where three isomers are quantified as two peaks when using capillary column chromatography, the calibration range of each peak should be considered separately, e.g., a diluted analysis is not required for total xylenes unless the concentration of either peak separately exceeds  $200 \mu g/L$ .

- D This flag identifies all compounds identified in an analysis at a secondary dilution factor. If a sample or extract is re-analyzed at a higher dilution factor, as in the "E" flag above, the "DL" suffix is appended to the sample number on the Form I for the diluted sample, and all concentration and quantification limit values reported on that Form I are flagged with the "D" flag.
- A This flag indicates that a TIC is a suspected aldol-condensation product.
- S This optional flag indicates that the compound is a matrix spike and thus, the concentration is not reported on Form I.
- P This flag is used for a pesticide/PCBs target analyte when there is greater than 25% difference for detected concentrations between the two GC Columns (See Form X). The lower of the two values is reported on the Form I and flagged with a "P".
- M This flag indicates that the analyte was quantified using a calibration curve constructed using a first or higher order regression fit as specified in and allowed by SW-846 methods 8260A (paragraph 7.3.6.1) and 8270B (paragraph 7.3.5.1).
- H The extraction holding time was exceeded.
- X, Y, or Z Other specific flags may be required to properly define the results. If used, they must be fully described and such description attached to the Sample Data Summary Package and the Sample Delivery Group (SDG) Narrative. For the data contained in this report, "Y" indicates that the data is to be used for qualitative purposes only and "Z" indicates that the initial calibration contains one less point than required by the method.

The combination of flags "BU" or "UB" is expressly prohibited. Blank contaminants are flagged "B" only when they are also detected in the sample.

For the analyses, other than volatile and semi-volatile organic compounds the lab qualifiers are as follows:

- U Used if the reported value is less than the instrument's detection limit (IDL).
- B Used if the reported value is less than ten times the IDL, but is greater than or equal to the IDL.
- N Used when spike recovery of either the matrix spike or matrix spike duplicate is not within the limits of 80 120%.
- E Used when the serial dilution or analytical spike is not within the SOW-156 limits.

# APPENDIX D DCS DATA

Table D-1. Evaporator parameters.

EVAPORATOR PARAMETERS, 0060STRT-1

Evapo	orator Paran	OR PARAME neters	1 ERS, 0060	51KI-1											FGAS PARAM	IETERS							_					
											e	D-150-1 vaporator	L-150-1 evaporator	F350-1 evaporator														
Time		T-150-2 degrees C			T-150-5 degrees C		T-150-7 degrees C	T-150-8 degrees C	T-150-9 degrees C	T-150-10 degrees C	ave	density ms/ml ir	level nches II	steam flow o/hour	T-336-1C degrees F	F 136-1 I	P-122-1 "wcvac	P-130-2 "wcvac	PD-130-1-1 IN WC	PD-130-2-1 IN WC	PD-130-3-1 IN WC	PD-130-4-1 IN WC	PD-130-1 IN WC	T-335-2 1 degrees F	<b>F-130-1-1</b> degrees F	T-130-2-1 degrees F	T-130-3-1 T degrees F	-130-4-1 F-130-1 degrees F scfm
05 JUN 01 09:30:00	70.274	72.117	75.015	75.114	46.441	45.187	36.15	36.742	37,146	37.729	53	1.1875	18.034	-2.387	75.648	170.52	11.581	48.987	0.008	-0.05€	3	0 1.9695	8.1261	183.11	105.49	104.86	98.824	163.65 632.45
05 JUN 01 09:45:00 05 JUN 01 10:00:00	96.436 97.442	96.992 98.077	96.509 97.413	96,394 97,443	47.203 69.385	45.97 69.732	36.431 37.98	37.083 38.581	37.547 37.948	38.185 38.702		1.1867 1.1858	37.086 56.134	-2.169 -1.95	75.623 75.662	170.05 169.59	11.582 11.583		0.008 0.008	-0.056 -0.056		0 1.9699 0 1.9703	8.1282 8.1304	183.09 183.09	105.49 105.49	104.86 104.85	98.82 98.816	163.64 606.83 163.64 634.13
05 JUN 01 10:15:00	98.377	99.135	98.247	98.44	89.207	89.009	42.735	43.22	39.106	40.699		1.185	75.182	-1.732	75.701	169.12	11.583	48.978	0.008	-0.056	3	0 1.9707	8.1326	183.08	105.48	104.85	98.812	163.64 629.7
05 JUN 01 10:30:00 05 JUN 01 10:45:00	99.023 99.273	99.529 99.77	99.081 99.393	99.175 99.433	94.15 94.931	93.396 94.915	50.974 64.089	51.409 64.633	45.414 58.734	47.101 60.355		1.1842 1.1833	93.734 111.05	-1.514 -1.295	75.74 75.779	168.66 168.19	11.584 11.585	48.974 49.971	800.0 800.0			0 1.9711 0 1.9715	8.1348 8.1369	183.07 183.06	105.48 105.48	104.84 104.83	98.808 98.804	163.64 630.73 163.64 633.6
05 JUN 01 11:00:00 05 JUN 01 11:15:00	99.524 99.775	100.01 100.25	99.644 99.896	99.691 99.949	94.538 93.141	95.159 93.095	81.304 91.544	82.062 92.22	77.126 90.363	78.993 92.638		1.1825 1.1817	127.73 137.23	-1.077 -0.859	75.818 75.857	167.73 167.26	11.586 11.587	48.968 48.965	0.008 800.0			0 1.9719 0 1.9722	8.1391 8.1413	183.05 183.04	105.48 105.48	104.83 104.82	98.8 98.796	163.63 647.47 163.63 625.8
05 JUN 01 11:30:00	99.987	100.25	100.23	100.4	92.836	92.706	96.146	96.955	96.523	98.08		1.1809	137.23	7.1828	75.896	166.8	11.587	48.961	0.008			0 1.9726		183.03	105.46	104.82	98.792	163.63 651.66
05 JUN 01 11:45:00 05 JUN 01 12:00:00	100.19 100.39	100.64 100.83	100.8 101.37	100.9 101.39	93.912 94.95	93.801 94.849	98.103 99.334	98.901 100.14	98.436 99.849	100.03 101.2		1.18 1.1792	137.02 136.91	691.63 1125.7	75.935 75.975	166.33 165.87	11.588 11.589	48.958 48.955	800.0 800.0	-0.056 -0.056		0 1.973 0 1.9734	8.1456 8.1478	183.02 183.01	105.47 105.47	104.81 104.8	98.788 98.784	163.63 646.97 163.63 614.2
05 JUN 01 12:15:00	100.59	101.02	101.85	101.89	95.988	95.897	100.04	100.79	100.51	101.77	400	1.1752	136.81	1652.5	76.014	165.4	11.59	48.952	0.008	-0.056	i	0 1.9738	8.15	183.00	105.47	104.8	98.78	163.62 627.16
05 JUN 01 12:30:00	100.79	101.22	101.97	102.01	97.025	96.945	99.686	100.38	100.23	101.85	100	1.1675	136.7	1725.5	76.053	164.93	11.591	48.949	0.008	-0.056	•	0 1.9742	8.1522	182.99	105.47	104.79	98.776	163.62 636.2
	EVAPORAT	OR PARAME	TERS, 0060	END-1																								
												D-150-1 vaporator	L-150-1 evaporator	F350-1 evaporator									_					
Time				T-150-4 degrees C			T-150-7 degrees C	T-150-8 degrees C		T-150-10 degrees C	Gra	density ms/ml in	level	steam flow hour	T-336-1C degrees F	F 136-1 F	P-122-1 "wcvac	P-130-2 "wcvac	PD-130-1-1 IN WC	PD-130-2-1 IN WC	PD-130-3-1 IN WC	PD-130-4-1 IN WC	PD-130-1 IN WC		f-130-1-1 degrees F	T-130-2-1 'degrees F	T-130-3-1 T degrees F	-130-4-1 F-130-1 degrees F scfm
05 JUN 01 16:00:00	103.58	103.92	104.57	103.98	97.792	97.612	101.25	102.32	101.33	102.7	102	1.2091	134.12	1719.5	76.599	158.42	11.602	48.903	0.008	-0.056		0 1.9796	8.1826	182.84	105.44	104.71	98.721	163.59 619.49
05 JUN 01 16:15:00 05 JUN 01 16:30:00	103.77 103.9	104.11 104.29	104.72 104.87	104.17 104.36	98.301 98.858	98.126 98.641	101.4 101.51	102.49 102.66	101.61 102.01	102.95 103.1		1.2129 1.2167	134.78 135.33	1744 1725.4	76.638 76.678	157.95 157.49	11.602 11.603	48.9 48.897	800.0 800.0	-0.056 -0.056		0 1.98 0 1.9804	8.1848 8.1869	182.83 182.82	105.44 105.43	104.7 104.7	98.717 98.714	163.59 603.97 163.59 630.28
05 JUN 01 16:45:00	104.03	104.41	105.01	104.55 104.74	99.415 99.778	99.155 99.555	101.61 101.72	102.83 103	102.41 102.82	103.25 103.4		1.2205	135.59 135.86	1732.1 1734.6	76.717 76.756	157.02 156.56	11.604 11.605	48.894 48.891	0.008 800.0	-0.056 -0.056		0 1.9807 0 1.9811	8.1891 8.1913	182.81 182.8	105.43	104.69	98.71	163.59 622.32 163.59 627.61
05 JUN 01 17:00:00 05 JUN 01 17:15:00	104.17 104.3	104.54 104.67	105.16 105.31	104.74	100.06	99.722	101.72	103.12	103.22	103.55		1.2242 1.228	135.99	1729.7	76.795	156.09	11.606	48.887	0.008	-0.056		0 1.9815	8.1935	182.79	105.43 105.43	104.69 104.68	98.706 98.702	163.58 610.26
05 JUN 01 17:30:00 05 JUN 01 17:45:00		104.79 104.92	105.46 105.61	105.01 105.1	100.34 100.63	99.889 100.06	101.93 102.03	103.2 103.28	103.62 103.64	103.7 103.85		1.2318 1.2356	135.83 135.67	1721.3 1729.2	76.834 76.873	155.26 154.41	11.606 11.607	48.884 48.881	0.008 0.008	-0.056 -0.056		0 1.9819 0 1.9823	8.1956 8.1978	182.78 182.77	105.43 105.43	104.67 104.67	98.698 98.694	163.58 642.05 163.58 626.59
05 JUN 01 18:00:00	104.69	105.05	105.76	105.19	101.04	100.22	102.14	103.35	103.23	104		1.2394	135.52	1714.8	76.912	153.55	11.608	48.878	0.008	-0.056		0 1.9827	8.2	182.76	105.42	104.66	98.69	163.58 650.26
05 JUN 01 18:15:00 05 JUN 01 18:30:00	104.82 104.95	105.17 105.3	105.83 105.81	105.28 105.37	101.46 101.87	100.39 100.74	102.25 103.18	103.43 103.51	103.06 103.68	104.15 105.07		1.2431 1.2469	135.36 135.56	1729.4 1725.5	76.951 76.99	152.69 151.83	11.609 11.61	48.874 48.871	0.008 0.008	-0.056 -0.056		0 1.9831 0 1.9835	8.2022 8.2043	182.75 182.74	105.42 105.42	104.66 104.65	98.686 98.682	163.58 625.62 163.57 618.58
05 JUN 01 18:45:00	105.08	105.43	105.8	105.46	102.29	101.12	103.21	103.59	103.72	105.12		1.2507	136.01	1730.3	77.029	150.98	11.61	48.868	0.008	-0.056		0 1.9838	8.2065	182.73	105.42	104.65	98.678	163.57 604.61
05 JUN 01 19:00:00 05 JUN 01 19:15:00	105.21 105.34	105.55 105.68	105.78 105.77	105.55 105.64	102.7 103.11	101.5 101.88	103.07 102.93	103.66 103.74	103.57 103.42	104.97 104.82		1.2545 1.2583	135.88 135.75	1714.2 1728.1	77.068 77.107	150.12 149.26	11.611 11.612	48.865 48.862	0.008 0.008	-0.056 -0.056		0 1.9842 0 1.9846	8.2087 8.2109	182.72 182.71	105.42 105.41	104.64 104.63	98.674 98.67	163.57 613.46 163.57 615.79
05 JUN 01 19:30:00 05 JUN 01 19:45:00	105.47 100.64	105.81 101.46	105.75 101.48	105.73 101.16	103.3 102.86	102.69 103.02	102.79 102.65	103.82 103.89	103.27 103.12	104.66 104.51	104	1.262 1.2658	135.62 135.49	1714.5 1726.9	77.146 77.185	148.4 147.55	11.613 11.613	48.858 48.855	0.008 0.008	-0.056 -0.056		0 1.985 0 1.9854	8.213 8.2152	182.7 182.69	105.41 105.41	104.63 104.62	98.666 98.662	163.57 649.86 163.56 626.28
00 0011 01 10.40.00		OR PARAME			102.00	,00.02	102.00		700.72	101.01			100.10		771.00	147.00	11,010	,0.000	0.000	0.000		7.5554	0.2102	102.00	100.41	104.02	00.002	100.00
	2777 01011	01111111111	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	<u> </u>								D-150-1 vaporator	L-150-1 evaporator	F350-1 evaporator									_					
Time							T-150-7	T-150-8		T-150-10		density	level	steam flow			_			PD-130-2-1	PD-130-3-1		PD-130-1		_	_		130-4-1 F-130-1
06 JUN 01 07:15:00	degrees C 61.829	degrees C 63.465	degrees C o 69.791	degrees C 70.188	degrees C 43.909	degrees C o 43.728	degrees C 36.42	degrees C 37.092	degrees C 37.571	degrees C 38.548	Gra	ms/ml In 1.1902	ches It 136.57	/hour 0.969	degrees F 76.543	scfm 168.77	"wcvac 11.553	"wcvac 48.707	IN WC 0.008	IN WC -0.056	IN WC	IN WC 0 2.0576	IN WC 8.3564	degrees F 183.03	degrees F 104.61	degrees F 104.43	degrees F 97.829	degrees F scfm 163.41 611.81
06 JUN 01 07:30:00	61.688	62.775 63.93	70.231 71.572	70.523 71.408	44.131 44.353	43.903 44.078	36.469 36.519	37.142 37.191	37.769 38.006	38.676 38.943		1.1896 1.1891	136.52 136.46	-0.588 -0.208	76.558 76.572	169 169.24	11.551 11.549	48.704 48.7	0.008 800.0	-0.056 -0.056		0 2.0575 0 2.0574	8.3562	183.03	104.61 104.61	104.43 104.43	97.833	163.41 635.14
06 JUN 01 07:45:00 06 JUN 01 08:00:00	62.838 96.666	97.526	97.21	96.776	44.575	44.445	36.568	37.191	38.243	39.21		1.1885	136.4	3.6601	76.586	169.47	11.548	48.7	0.008	-0.056		0 2.0573	8.356 8.3559	183.02 183.02	104.61	104.43	97.836 97.84	163.41 639.53 163.4 650.84
06 JUN 01 08:15:00 06 JUN 01 08:30:00	97.881 98.485	98.537 99.254	98.028 98.592	98.157 99.204	59.743 82.048	59.993 82.078	37.463 40.546	38.102 41.242	38.48 38.718	39.477 39.745		1.188 1.1875	136.35 136.33	446.62 871.3	76.6 76.614	169.71 169.94	11.546 11.544	48.7 48.7	0.008 0.008	-0.056 -0.056		0 2.0572 0 2.0571	8.3557 8.3556	183.02 183.01	104.62 104.62	104.44 104.45	97.844 97.848	163.4 666.56 163.4 633.06
06 JUN 01 08:45:00	99.041	99.972	99.156	99.457	93.676	92.752	47.382	47.542	41.907	43.421		1.1793	136.66	1335.8	76.629	170.18	11.543	48.7	0.008	-0.056		0 2.0569	8.3554	183.01	104.62	104.45	97.851	163.4 615.81
06 JUN 01 09:00:00 06 JUN 01 09:15:00	99.559 99.893	100.3 100.55	99.72 100.18	99.71 99.964	96.311 95.898	95.872 95.598	57.503 73.257	58.15 73.735	51.775 67.856	53.565 69.073		1.1735 1.1683	136.42 136.06	1736.2 1735.3	76.643 76.657	170.42 170.65	11.541 11.539	48.7 48.7	800.0 800.0	-0.056 -0.056		0 2.0568 0 2.0567	8.3553 8.3551	183 183	104.62 104.62	104.45 104.46	97.855 97.859	163.39 637.62 163.39 642.69
06 JUN 01 09:30:00	100.23	100.81	100.59	100.35	95.484	95.176	88.924	89.958	85.873	88.072		1.1631	135.7	1721.4	76.671 76.685	170.89	11.537	48.7 48.7	0.008	-0.056		0 2.0566	8.355	182.99	104.62	104.46	97.863	163.39 643.56
06 JUN 01 09:45:00 06 JUN 01 10:00:00	100.56 100.89	101.05 101.26	101 101.41	100.86 101.36	93.178 94.269	94.227 94.711	94.626 98.045	96.075 98.89	94.927 98.442	96.641 99.804		1.1628 1.1668	132.97 129.4	1724.7 1731	76.699	171.12 171.36	11.536 11.534	48.7	800.0 800.0	-0.056 -0.056		0 2.0565 0 2.0564	8.3548 8.3547	182.99 182.99	104.62 104.63	104.46 104.47	97.866 97.87	163.39 652.44 163.38 662.49
06 JUN 01 10:15:00 06 JUN 01 10:30:00	101.14 101.35	101.47 101.68	101.82 102.23	101.87 102.16	95.972 97.357	95.819 97.109	99.561 100.54	100.39 101.16	100.08 100.93	101.31 102.19		1.1714 1.1781	125.45 121.13	1726.2 1725.9	76.714 76.728	171.59 171.83	11.532 11.531	48.7 48.7	0.008 800.0	-0.056 -0.056		0 2.0563 0 2.0562	8.3545 8.3544	182.98 182.98	104.63 104.63	104.47 104.48	97.874 97.878	163.38 644.66 163.38 620.82
06 JUN 01 10:45:00	101.56	101.89	102.43	102.37	98.629	98.262	101.01	101.52	100.71	102.8		1.1848	116.85	1724.8	76.742	172.06	11.529	48.7	0.008	-0.056		0 2.0561	8.3542	182.97	104.63	104.48	97.881	163.38 630.82
06 JUN 01 11:00:00 06 JUN 01 11:15:00	101.77 101.98	102.1 102.31	102.56 102.69	102.58 102.77	98.601 95.504	98.668 95.169	101.05 97.614	100.93 98.603	100.25 98.546	102.24 99.762		1.1916 1.1963	113.01 123.25	1730 1724.7	76.756 76.77	172.3 172.53	11.527 11.525	48.7 48.7	0.008 0.008	-0.056 -0.056		0 2.056 0 2.0559	8,354 8,3539	182.97 182.97	104.63 104.63	104.48 104.49	97.885 97.889	163.37 664.12 163.37 634.06
06 JUN 01 11:30:00	102.19	102.52	102.82	102.94	94.835	95.309	98.235	99.465	99.119	100.27		1.1994	124.3	1725.3	76.785	172.77	11.524	48.7	0.008	-0.056		0 2.0558	8.3537	182.96	104.63	104.49	97.893	163.37 656.65
06 JUN 01 11:45:00	102.4	102.73	102.94	103.12	94.85	95.45	98.661	99.799	99.472	100.59		1.2025	125.62	1728.1	76.799	173	11.522	48.7	0.008	-0.056		0 2.0557	8.3536		104.63	104.49	97,897	163.37 664.84
	EVAPORAT	OR PARAME	IEKS, 0060	END-Z					•			D-150-1	L-150-1	F350-1									_					
Time		T-150-2				T-150-6			T-150-9			vaporator density	evaporator level			F 136-1 P					PD-130-3-1							130-4-1 F-130-1
06 JUN 01 15:00:00	degrees C 104.59	degrees C 105.44	degrees C of 105.51	degrees C o 105.41	degrees C 99.217	degrees C of 98.681	degrees C 102.98	degrees C 103.27	degrees C 103.11	degrees C 104.13	Gra	ms/ml In 1.2536	ches lb 128.9	/hour 1732.2	degrees F 76.983	scfm 176.06	"wcvac 11.5	"wcvac 48.7	IN WC 0.008	IN WC -0.056	IN WC	IN WC 2.0544	IN WC 8.3516	degrees F 182.91	degrees F 104.65	degrees F 104.54	degrees F 97.945	degrees F scfm 163.33 661.48
06 JUN 01 15:15:00	104.76	105.54	105.63	105.59	99.421	98.844	103.12	103.53	103.33	104.4		1.257	129.23	1725.5	76.997	176.3	11.498	48.7	0.008	-0.056		2.0543	8.3514	182.9	104.65	104.55	97.949	163.33 641.36
06 JUN 01 15:30:00 06 JUN 01 15:45:00	104.92 105.09	105.65 105.76	105.67 105.67	105.77 105.94	99.626 99.83	99.007 99.17	103.27 103.41	103.79 104.04	103.49 103.66	104.67 104.93		1.2604 1.2638	129.47 129.67	1716.1 1722.5	77.011 77.026	176.54 176.77	11.496 11.495	48.7 48.7	0.008 0.008	-0.056 -0.056		2.0542 2.0541	8.3513 8.3511	182.9 182.9	104.65 104.65	104.55 104.55	97.953 97.957	163.33 620.13 163.33 660.25
06 JUN 01 16:00:00 06 JUN 01 16:15:00	105.25 105.42	105.87 105.98	105.67 105.67	106.12 106.29	100.04 100.24	99.333 99.497	103.55 103.69	104.3 104.56	103.83 103.99	105.2 105.47		1.2672 1.2706	129.86 130.06	1724.8 1726.5	77.04 77.054	177.01 177.24	11.493 11.491	48.7 48.7	800.0 800.0	-0.056 -0.056	(	2.054	8.351 8.3508	182.89 182.89	104.66 104.66	104.56 104.56	97.96 97.964	163.32 667.6 163.32 649.2
06 JUN 01 16:30:00	105.58	106.09	105.67	106.47	100.44	99.66	103.73	104.82	104.16	105.74		1.274	130.25	1727.6	77.068	177.48	11.49	48.7	0.008	-0.056	(	2.0538	8.3507	182.88	104.66	104.57	97.968	163.32 652.49
06 JUN 01 16:45:00 06 JUN 01 17:00:00	105.75 105.91	106.19 106.3	105.67 105.67	106.65 106.82	100.65 100.85	99.823 99.986	103.71 103.68	104.95 104.79	104.32 104.49	106.01 106.27		1.2775 1.2809	130.43 130.03	1733.7 1731.7	77.082 77.097	177.71 177.95	11.488 11.486	48.7 48.7	0.008 0.008	-0.056 -0.056	(	2.0537 2.0536	8.3505 8,3504	182.88 182.88	104.66 104.66	104.57 104.57	97.972 97.975	163.32 642.31 163.31 656.42
06 JUN 01 17:15:00	106.08	106.41	105.67	107	101.06	100.15	103.65	104.62	104.65	106.54		1.2843	129.64	380.21	77.111	178.18	11.484	48.7	0.008	-0.056		2.0535	8.3502	182.87	104.66	104.58	97.979	163.31 626.64
06 JUN 01 17:30:00 06 JUN 01 17:45:00	101.12 99.677	101.87 100.23	102 100.5	101.54 99.811	100.8 100.48	100.31 99.884	103.62 103.59	104.46 104.29	104.53 104.01	106.04 105.32		1.2877 1.2911	129.31 128.98	-9.053 -8.939	77.125 77.139	178.42 178.65	11.483 11.481	48.7 48.7	0.008 0.008	-0.056 -0.056	(	2.0534 2.0533	8.3501 8.3499	182.87 182.86	104.66 104.66	104.58 104.58	97.983 97.987	163.31 668.24 163.31 634.91
06 JUN 01 18:00:00	97.996	99.212	99.361	98.171	100.15	99.339	103.29	104.13	103.32	104.49		1.2938	128.65	-8.824	77.153	178.89	11.479	48.7	0.008	-0.056	(	2.0532	8.3497	182.86	104.67	104.59	97.99	163.3 658.67

Table D-1. Evaporator parameters.

EVAPORATOR PARAMETERS, 0050STRT-1

	•		STRT-1							D-150-1	L-150-1	F350-1									_						
Time T-150	0-1 T-150-2	T-150-3	T-150-4 T	150-5 T-1	50-6 T-	150-7 T-	150-8	T-150-9	T-150-10	evaporator density	evaporator level	evaporator steam flow	T-336-1C	F 136-1 P	122-1 P	-130-2 F	PD-130-1-1	PD-130-2-1	PD-130-3-1	PD-130-4-1	PD-130-1 _		-130-1-1 T	-130-2-1 T	-130-3-1 T-	130-4-1 F-130	0-1
07 JUN 01 08:00:00 96.9	es C degrees C i.912 97.913 '.748 98.768	96.931 98.67	degrees C de 96.902 97.801	8.512	48.563	grees C deg 41.697 42.467	grees C d 42.07 43.437	egrees C 41.993 42.053	degrees C 42.81 42.887	Grams/ml 1.1742 1.1737	136.8 136.74	lb/hour 4.2814 678.65	degrees F 77.948 77.962	scfm 171.82 170.91	"wcvac 11.384 11.382	"wcvac 48.689 48.69	IN WC 0.008 0.008	IN WC -0.056 -0.056	IN WC	IN WC 2.0474 2.0473	IN WC 8.3412 8,341	degrees F 182.63 182.63	degrees F 104.74 104.74	degrees F 104.8 104.8	degrees F 98.201 98.204	163.16 6	cfm 655.98 676.63
07 JUN 01 08:30:00 98.4	3.444 99.431 3.139 99.984	99.194 99.643	98.593 99.386	92.052		47.424 55.84	48.118 56.547	43.667 50.83	45.39 52.693	1.1732 1.1716	136.67 136.68	1095.9 1522,7	77.976 77.99	170 169.09	11.38 11.379	48.692 48.693	0.008 0.008	-0.056 -0.056	Č	2.0472	8.3408 8.3407	182.62 182.62	104.74 104.74	104.81 104.81	98.208 98.212	163.16 6	668.63 629.8
	.561 100.23	99.876 100.11	99.768 100.08	95.943		69.314 86.41	70.263 87.072	65.202 82.498	66.09 84.708	1.1606 1.1542	137.23 136.81	1715.6 1723.3	78.004 78.018	168.17 167.26	11.377 11.375	48.695 48.696	0.008 0.008	-0.056 -0.056	C	2.047	8.3405 8.3404	182.61 182.61	104.74 104.74 104.74	104.81 104.82	98.216 98.219	163.15	663.7 626.7
07 JUN 01 09:30:00 100	0.19 100.74	100.11 100.34 100.56	100.39	93.773	93.773		94.365 98.067	93.391 97.605	95.257 98.99	1.1574 1.1634	134.05	1736	78.033	166.35	11.373	48.698	0.008	-0.056	0	2.0468	8.3402	182.61	104.74	104.82	98.223	163.15 6	656.16 652.55
07 JUN 01 10:00:00 100	0.64 101.12	100.75	100.85	94.819	94.618	98.815	99.615	99.357	100.69	1.1674	130,55 126,72	1729.2 1729.6	78.047 78.061	165.44 164.54	11.372	48.699 48.701	0.008 0.008	-0.056 -0.056	C	2.0467	8.3401 8.3399	182.6 182.6	104.74 104.75	104.82 104.83	98.227 98.231	163.14 6	628.17
07 JUN 01 10:30:00 101	0.85 101.3 1.07 101.48	100.94 101.14	101.07 101.3	96.983	96.531	99.789 100.34	100.57	100.39	101.61 102.09	1.1716 1.1758	122.42 118.31	1730 1730	78.075 78.089	163.63 162.72	11.368 11.367	48.702 48.704	0.008 0.008	-0.056 -0.056	0	2.0464	8.3398 8.3396	182.59 182.59	104.75 104.75	104.83 104.84	98.234 98.238	163.14 6	651.21 619.94
	1.28 101.66 01.5 101.84	101.33 101.52	101.53 101.75			99.844 97.425	100.63 98.163	99.932 97.966	102.49 99.678	1.1798 1.1837	114.36 122.13	1732.5 1730.6	78.103 78.117	161.81 161.61	11.365 11.363	48.706 48.707	0.008 0.008	-0.056 -0.056	0	2.0463 2.0462	8.3394 8.3393	182.59 182.58	104.75 104.75	104.84 104.84	98.242 98.246		641.03 645.22
EVAPO	ORATOR PARAM	ETERS, 0050	END-1							D-150-1	L-150-1	F350-1									_						
Time T-150	0-1 T-150-2	T-150-3	T-150-4 T	150-5 T-1	50-6 T-	150-7 T-	150-8 °	T-150-9	T-150-10	evaporator density	evaporator level	evaporator	T-336-1C	F 136-1 P-	122-1 P	130-2 F	PD-130-1-1	PD-130-2-1	PD-130-3-1	PD-130-4-1	PD-130-1	T-335-2 T	-130-1-1 T	-130-2-1 T	-130-3-1 T-	130-4-1 F-130	0-1
			degrees C de 103.67				rees C de 101.38	egrees C 100.95	degrees C 102.44	Grams/ml 1.23	Inches 132.43	lb/hour 1731.7	degrees F 78.288	scfm 163.34	"wcvac 11.343	"wcvac 48,725	IN WC 0.008	IN WC -0.056	IN WC	IN WC 2.045	_		degrees F 104.77	degrees F 104.89		degrees F sc	cfm 617.59
	3.42 103.96 3.55 104.09	104.02 104.22	103.79 103.94		97.443 97.973	100.92 101.3	101.72 102.07	101.21 101.64	102.67 103.12	1.2339 1.2377	133.71 134.6	1731.9 1736.1	78.302 78.316	163.49 163.63	11.341 11.339	48.727 48.729	0.008 0.008	-0.056 -0.056	0	2.0449 2.0448	8.3373 8.3371	182.53 182.52	104.77 104.77	104.89 104.9	98.294 98.298		668.63 637.74
	3.62 104.22 3.68 104.34	104.41 104.6	104.12 104.3			101.67 101.92	102.42 102.61	102.07 102.35	103.56 103.86	1.2416 1.2454	134.9 133.81	1725.1 1732.4	78.33 78.344	163.77 163.92	11.338 11.336	48.73 48.732	0.008 0.008	-0.056 -0.056	0	2.0447 2.0446	8.337 8.3368	182.52 182.52	104.77 104.77	104.9 104.9	98.302 98.306		640.39 645.49
07 JUN 01 15:15:00 103	3.74 104.47 3.79 104.6	104.79 104.99	104.48 104.65	99.751	99.52	102.08 102.23	102.76 102.91	102.53 102.7	104.05 104.24	1.2493 1.2531	132.47 131.81	1745.4 1726.2	78.358 78.373	164.06 164.21	11.334 11.333	48.733 48.735	0.008 0.008	-0.056 -0.056	0	2.0445 2.0444	8.3367 8.3365	182.51 182.51	104.77 104.77	104.91 104.91	98.31 98.313	163.09 6	644.63 616.02
	3.85 104.72	105.18 105.29	104.83 105.01	100.16	99.857	102.39 102.51	103.06 103.21	102.88 102.95	104.43 104.54	1.257 1.2609	132.18 133.08	1728.6 1730.2	78.387 78.401	164.35 164.5	11.331 11.329	48.736 48.738	0.008 0.008	-0.056 -0.056	0	2.0442 2.0441	8.3364 8.3362	182.5 182.5	104.77 104.78	104.91 104.92	98.317 98.321	163.08 6	623.87 631.34
07 JUN 01 16:15:00 103	3.96 104.98 4.02 105.1	105.36 105.42	105.18 105.36	100.52	100.19	102.53 102.55	103.26 103.27	103 103.04	104.59 104.65	1.2647 1.2686	134.24 135.06	1709 1725.9	78.415 78.429	164.64 164.78	11.327 11.326	48.739 48.741	0.008 0.008	-0.056 -0.056	0	2.044 2.0439	8.3361 8.3359	182.5 182.49	104.78 104.78	104.92 104.93	98.325 98.328	163.08 6	654.16 635.04
07 JUN 01 16:45:00 104	4.07 105.23 4.13 105.36	105.49 105.56	105.54 105.72	101.05		102.58 102.6	103.28	103.08 103.13	104.71 104.77	1.2724 1.2763	134.52 134.84	1729.7 1727.4	78.444 78.458	164.93 165.07	11.324 11.322	48.742 48.744	0.008	-0.056 -0.056	0	2.0438 2.0437	8.3358 8.3356	182.49 182.48	104.78 104.78	104.93 104.93	98.332 98.336	163.07 6	643.47 644.53
	4.19 105.48	105.62	105.89	101.88	101.47	102.62	103.31	103.17	104.83	1.2802	135.36	1723.2	78.472	165.22	11.321	48.745	0.008	-0.056	0	2.0436	8.3355	182.48	104.78	104.94	98.34		651.51
EVAPO	ORATOR PARAM	ETERS, 00505	STRT-2					<del></del>		D-150-1	L-150-1	F350-1									_						
Time T-150-	0-1 T-150-2	T-150-3	T-150-4 T-	150-5 T-1	50-6 T-	150-7 T-	150-8 1	T-150-9	T-150-10	evaporator density	evaporator ievel	evaporator steam flow	T-336-1C	F 136-1 P-	122-1 P-	130-2 P	PD-130-1-1	PD-130-2-1	PD-130-3-1	PD-130-4-1	PD-130-1	T 225 2 T			120 2 1 T	130-4-1 F-130	0-1
4				<del></del>					1		La ala a a	11. 4.		<del></del>													
11 JUN 01 07:15:00 69.9	.962 71.629	degrees C o 72.958	degrees C deg 73.277	rees C degr 45.649 4	ees C deg 44.603	rees C deg 41.619	rees C de 41.541	41.857	degrees C 42.859	Grams/ml 1.1754	137.02	lb/hour -2.732		56.42 11	"wcvac 8.0286	"wcvac 48.7	IN WC 0.008	IN WC -0.056	IN WC	IN WC 2.0152	IN WC 8.4546	degrees F 183.33	degrees F 105.66	degrees F 105.56	degrees F 98.865	degrees F sc 162.98 6	cfm 635.69
11 JUN 01 07:15:00 69.9 11 JUN 01 07:30:00 67.4 11 JUN 01 07:45:00 65.4	.962 71.629 .499 69.081 .488 67.129	degrees C o 72.958 70.876 69.332	73.277 71.367 69.758	rees C degr 45.649 4 45.126 4 45.428 4	ees C deg 14.603 4 14.854 4 15.105 4	rees C deg 41.619 41.663 41.706	rees C de 41.541 41.587 41.634	41.857 41.897 41.937	42.859 42.899 42.94	Grams/ml 1.1754 1.175 1.1746	137.02 136.99 136.97	-2.732 -2.66 -2.587	74.848 1 74.825 74.802	56.42 11 156.5 156.57	8.0286 8.0294 8.0302	48.7 48.7 48.7	0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056		IN WC 2.0152 2.0151 2.0151	IN WC 8.4546 8.4571 8.4597	degrees F 183.33 183.33 183.33	degrees F 105.66 105.65 105.65	degrees F 105.56 105.56 105.56	degrees F 98.865 98.866 98.867	degrees F sc 162.98 6: 162.97 6 162.96 6-	cfm 635.69 614.51 649.32
11 JUN 01 07:15:00 69.9 11 JUN 01 07:30:00 67.4 11 JUN 01 07:45:00 65.4 11 JUN 01 08:00:00 63.8 11 JUN 01 08:15:00 97.0	.962 71.629 .499 69.081 .488 67.129 .843 65.437 .019 97.79	degrees C 72.958 70.876 69.332 68.129 97.328	res C degrees C	rees C degrees C	ees C deg 44.603 4 44.854 4 45.105 4 45.355 51.182 4	rees C deg 41.619 41.663 41.706 41.75 41.793	rees C de 41.541 41.587 41.634 41.681 41.727	41.857 41.897 41.937 41.977 42.017	42.859 42.899 42.94 42.98 43.02	Grams/ml 1.1754 1.175 1.1746 1.1742 1.1737	137.02 136.99 136.97 136.94 136.92	-2.732 -2.66 -2.587 -2.514 4.5011	74.848 1 74.825 74.802 74.78 74.757	56.42 11 156.5 156.57 156.65 156.73	8.0286 8.0294 8.0302 8.0309 8.0317	48.7 48.7 48.7 48.7 48.7	0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056		IN WC 2.0152 2.0151 2.0151 2.015 2.015	IN WC 8.4546 8.4571 8.4597 8.4622 8.4647	degrees F 183.33 183.33 183.33 183.33	degrees F 105.66 105.65 105.65 105.65 105.65	degrees F 105.56 105.56 105.56 105.56 105.56	degrees F 98.865 98.866 98.867 98.869 98.87	degrees F sci 162.98 6: 162.97 6 162.96 6: 162.96 6: 162.95 6:	cfm 635.69 614.51 649.32 601.31 652.47
11 JUN 01 07:15:00 69.9 11 JUN 01 07:30:00 67.4 11 JUN 01 07:45:00 65.4 11 JUN 01 08:00:00 63.8 11 JUN 01 08:15:00 97.0 11 JUN 01 08:30:00 97.8 11 JUN 01 08:45:00 94.1	.962 71.629 .499 69.081 .488 67.129 .843 65.437 .019 97.79 .852 98.832 .106 95.214	degrees C 72.958 70.876 69.332 68.129 97.328 98.004 94.831	regrees C degrees C degrees C 73.277 71.367 69.758 68.417 97.059 98.247 94.656	rees C degri 45.649 4 45.126 4 45.428 4 45.729 4 50.646 5 81.727 8 83.274 8	ees C deg 44.603 4 44.854 4 45.105 4 45.355 51.182 4 31.871 4 32.282 4	rees C deg 41.619 41.663 41.706 41.75 41.793 42.292 47.058	41.541 41.587 41.634 41.681 41.727 42.943 47.149	41.857 41.897 41.937 41.977 42.017 42.057 42.285	42.859 42.899 42.94 42.98 43.02 43.06 43.323	Grams/ml 1.1754 1.175 1.1746 1.1742 1.1733 1.1729	137.02 136.99 136.97 136.94 136.92 136.89 136.87	-2.732 -2.66 -2.587 -2.514 4.5011 434.46 -11.47	74.848 1 74.825 74.802 74.78 74.757 74.735 74.712	56.42 11 156.5 156.57 156.65 156.73 156.8 156.88	8.0286 8.0294 8.0302 8.0309 8.0317 8.0325 8.0333	48.7 48.7 48.7 48.7 48.7 48.7 48.7	0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056 -0.056		IN WC 2.0152 2.0151 2.0151 2.015 2.015 2.015 2.0149	8.4546 8.4571 8.4597 8.4622 8.4647 8.4673 8.4698	degrees F 183.33 183.33 183.33 183.33 183.32 183.32 183.32	degrees F 105.66 105.65 105.65 105.65 105.65 105.65	degrees F 105.56 105.56 105.56 105.56 105.56 105.56 105.56	degrees F 98.865 98.866 98.867 98.869 98.87 98.871 98.873	degrees F sci 162.98 6: 162.97 6 162.96 6: 162.96 6: 162.95 6: 162.94 6:	cfm 635.69 614.51 649.32 601.31 652.47 634.15 627.71
11 JUN 01 07:15:00 69.9 11 JUN 01 07:30:00 67.4 11 JUN 01 07:45:00 65.4 11 JUN 01 08:00:00 63.8 11 JUN 01 08:30:00 97.0 11 JUN 01 08:45:00 94.1 11 JUN 01 09:15:00 98.2	.962 71.629 .499 69.081 .488 67.129 .843 65.437 .019 97.79 .852 98.832 .106 95.214 .396 93.392 .239 98.739	degrees C 72.958 70.876 69.332 68.129 97.328 98.004 94.831 92.62 98.249	regrees C degrees C 73.277 71.367 69.758 68.417 97.059 98.247 94.656 92.682 98.403	rees C degri 45.649 4 45.126 4 45.428 4 45.729 4 50.646 5 81.727 8 83.274 8 70.97 6 89.366 8	ees C deg 44.603 4 44.854 4 45.105 4 45.355 51.182 4 31.871 4 32.282 4 58.946 5 38.584 5	rees C deg 41.619 41.663 41.706 41.75 41.793 42.292 47.058 50.947 52.842	rees C de 41.541 41.587 41.634 41.681 41.727 42.943 47.149 51.152 53.489	41.857 41.897 41.937 41.977 42.017 42.057 42.285 42.861 49.053	42.859 42.899 42.94 42.98 43.02 43.06 43.323 43.894 50.57	Grams/ml 1.1754 1.1754 1.1756 1.1742 1.1737 1.1733 1.1729 1.1725 1.1711	137.02 136.99 136.97 136.94 136.92 136.89 136.87 136.84 136.84	-2.732 -2.66 -2.587 -2.514 4.5011 434.46 -11.47 -10.86 1556.5	74.848 1 74.825 74.802 74.78 74.757 74.735 74.712 74.689 74.71	56.42 11 156.5 156.57 156.65 156.73 156.8 156.88 156.77 156.64	8.0286 8.0294 8.0302 8.0309 8.0317 8.0325 8.0333 11.451 11.448	48.7 48.7 48.7 48.7 48.7 48.7 48.7 48.7	0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056		IN WC 2.0152 2.0151 2.0151 2.015 2.015 2.015 2.0149 2.0149 2.0148	8.4546 8.4571 8.4597 8.4622 8.4647 8.4673 8.4698 8.4666 8.4628	degrees F 183.33 183.33 183.33 183.33 183.32 183.32 183.32 183.32 183.32	degrees F 105.66 105.65 105.65 105.65 105.65 105.65 105.65 105.65	degrees F 105.56 105.56 105.56 105.56 105.56 105.56 105.56 105.56	degrees F 98.865 98.866 98.867 98.869 98.871 98.871 98.873 98.874 98.875	degrees F sci 162.98 6: 162.97 6: 162.96 6: 162.96 6: 162.95 6: 162.94 6: 162.94 6: 162.94 6: 162.93 6:	cfm 635.69 614.51 649.32 601.31 652.47 634.15 627.71 661.54 616.68
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.0.56 101.31 .0.82 101.45 1.07 101.59  DRATOR PARAM  D-1 T-150-2 as C degrees C .278 103.02 .287 103.12 .296 103.22 .306 103.32 .315 103.42 .306 103.32 .315 103.42 .324 103.52 .334 103.62 .343 103.72 .352 103.82 .365 103.82	T-150-3 degrees C 72.958 70.876 69.332 68.129 97.328 98.004 94.831 92.62 98.249 98.711 99.125 99.459 99.792 100.13 101.46 101.67  T-150-3 degrees C 104.04 104.26 104.48 104.71 104.92 105.02 105.02 105.12 105.22 105.32 105.42	range C dec 73.277 71.367 69.758 68.417 97.059 98.247 94.656 92.682 98.403 99.065 99.672 99.932 100.14 100.35 100.56 100.77 100.98 101.19 FIND-2 T-150-4 T-150	rees C degree 45.649 445.649 445.649 445.729 445.729 45.664 5.664	ees C deg 44.603 44.854 45.105 45.105 45.355 45.355 45.355 45.355 45.355 45.355 46.35 46.42 46.42 47.104 48.44 49.8142	rees C deg 41.619 41.663 41.766 41.776 41.793 42.292 47.058 50.947 52.842 62.168 78.357 90.481 95.004 97.689 98.745 99.371 99.347 97.007  150.7  T-rees C deg 100.07 100.26 100.46 100.65 100.84 101.04 101.23 101.42 101.62	rees C de 41.541 41.587 41.634 41.681 41.727 42.943 47.149 51.152 53.489 62.969 98.53 99.671 100.31 100.02 97.942 100.02 97.942 100.02 97.942	41.857 41.897 41.937 41.977 42.017 42.057 42.285 42.861 49.053 57.829 73.355 89.382 99.471 100.01 100.11 99.637 97.735 Pegrees C 101.4 101.62 101.84 102.06 102.28 102.5 102.61 102.37 102.13 101.88	42.859 42.899 42.94 42.98 43.02 43.06 43.323 43.894 50.57 59.538 74.991 91.101 97.397 99.511 100.6 101.25 101.72 101.25 98.961  T-150-10 degrees C 102.8 103.01 103.23 103.44 103.65 103.87 104.08	Grams/ml 1.1754 1.1754 1.1756 1.1746 1.1742 1.1737 1.1733 1.1729 1.1725 1.1711 1.1643 1.1607 1.1582 1.159 1.1633 1.1675 1.1715 1.1755 1.1794 1.1834  D-150-1 evaporator density Grams/ml 1.235 1.2469 1.2509 1.2549 1.2549	137.02 136.99 136.97 136.94 136.92 136.89 136.87 136.84 136.82 137.05 137.39 136.35 132.79 128.99 125.09 120.8 116.68 113.27 123.08  L-150-1 evaporator level inches 33.55 132.21 132.65 132.01 130.75 129.79 129.89 131.13 133.32	-2.732 -2.66 -2.587 -2.514 4.5011 434.46 -11.47 -10.86 1556.5 1722.1 1722.3 1733.4 1730.3 1734.7 1730.5 1731 1718 1713.3 1731  F350-1 evaporator steam flow b/hour  1738.9 1732.2 1727.5 1728.8 1726.9 1735.3 1724.9 1720.2 1729.2 1731.6	74.848 1 74.825 74.802 74.78 74.757 74.735 74.712 74.689 74.71 74.765 74.82 74.875 74.929 74.984 75.039 75.094 75.149 75.204 75.259  T-336-1C degrees F 75.972 76.082 76.192 76.192 76.247 76.302 76.356 76.411 76.466	56.42 11 156.5 156.57 156.65 156.73 156.88 156.88 156.77 156.64 156.51 156.39 156.26 155.87 155.87 155.87 155.75 155.62 155.36 P	8.0286 8.0294 8.0309 8.0309 8.0317 8.0325 11.445 11.442 11.436 11.433 11.431 11.425 11.422 11.419 P-Wcvac 11.38 11.371 11.365 11.374 11.362 11.359 11.359 11.359	48.7 48.7 48.7 48.7 48.7 48.7 48.7 48.7	0.008 0.008	IN WC -0.056	IN WC 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	IN WC 2.0152 2.0151 2.0151 2.0155 2.0151 2.015 2.0149 2.0148 2.0148 2.0147 2.0147 2.0147 2.0146 2.0148 2.0148 2.0148 2.0147 2.0147 2.0147 2.0147 2.0146 2.0148 2.0145 2.0145 2.0138 2.0138 2.0138 2.0137 2.0137 2.0137 2.0137 2.0136 2.0135	IN WC 8.4546 8.4571 8.4597 8.4622 8.4647 8.4673 8.4698 8.4666 8.4628 8.4551 8.4554 8.4412 8.4405 8.4368 8.4331 8.4294 8.4256 IN WC 8.3773 8.3736 8.3699 8.3662 8.3624 8.3555 8.3513 8.3476 8.3438	degrees F 183.33 183.33 183.33 183.32 183.32 183.32 183.32 183.32 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.37 183.37 183.27 183.27 183.27 183.27 183.27 183.27 183.27 183.27 183.27 183.27	degrees F 105.66 105.65 105.65 105.65 105.65 105.65 105.65 105.65 105.64 105.64 105.64 105.64 105.64 105.64 105.63 105.63	degrees F 105.56	degrees F 98.865 98.866 98.867 98.867 98.871 98.871 98.873 98.875 98.875 98.876 98.879 98.88 98.881 98.883 98.884 98.885 98.887 98.888	degrees F sci 162.98 6 162.96 6 162.96 6 162.96 6 162.95 66 162.94 66 162.94 66 162.93 66 162.94 66 162.93 66 162.93 66 162.93 66 162.92 6 162.91 66 162.75 66 162.76 66 162.75 66 162.75 66 162.75 66 162.75 66 162.75 66 162.75 66 162.77 66 162.75 66 162.77 66	ofm 635.69 614.51 649.32 601.31 652.47 634.15 627.71 661.54 661.68 647.07 652.57 608.78 638.46 643.52 6413.6 626.29 640.44 624.31 627.7
11 JUN 01 07:15:00 69.9 11 JUN 01 07:45:00 67.4 11 JUN 01 08:00:00 63.8 11 JUN 01 08:15:00 97.0 11 JUN 01 08:30:00 97.8 11 JUN 01 08:45:00 94.1 11 JUN 01 08:45:00 94.1 11 JUN 01 09:30:00 98.6 11 JUN 01 09:30:00 98.6 11 JUN 01 09:45:00 98.2 11 JUN 01 10:30:00 99.7 11 JUN 01 10:30:00 99.7 11 JUN 01 10:30:00 99.7 11 JUN 01 11:15:00 100. 11 JUN 01 11:15:00 100. 11 JUN 01 11:45:00 101. 11 JUN 01 11:45:00 101. 11 JUN 01 15:45:00 101. 11 JUN 01 15:45:00 102. 11 JUN 01 15:45:00 103. 11 JUN 01 16:45:00 103. 11 JUN 01 16:45:00 103. 11 JUN 01 16:30:00 103. 11 JUN 01 16:30:00 103. 11 JUN 01 17:45:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:50:00 103. 11 JUN 01 17:45:00 103. 11 JUN 01 17:45:00 103. 11 JUN 01 17:45:00 103. 11 JUN 01 17:45:00 103.	9.962 71.629 4.89 69.081 4.88 67.129 8.43 65.437 0.19 97.79 8.52 98.832 1.06 95.214 3.396 93.392 2.39 98.739 6.18 99.22 9.997 99.63 2.81 99.94 1.00.56 101.31 0.05 101.31 1.07 101.59  DRATOR PARAM  D-1 T-150-2 BS C degrees C 2.78 103.02 2.87 103.02 2.87 103.02 2.87 103.02 2.88 103.02 2.89 103.22 3.15 103.42 3.34 103.52 3.34 103.62 3.34 103.62 3.34 103.62 3.352 103.82 3.62 103.92 3.71 104.02	T-150-3  degrees C 72.958 70.876 69.332 68.129 97.328 98.004 94.831 92.62 98.249 98.711 99.125 99.459 99.792 100.13 101.46 101.67  T-150-3  degrees C 104.04 104.26 104.48 104.71 104.92 105.02 105.12 105.22 105.32	range C dec 73.277 71.367 69.758 68.417 97.059 98.247 94.656 92.682 98.403 99.065 99.368 99.368 99.368 100.14 100.35 100.56 100.77 100.98 101.19 FIND-2 T-150-4 T-16grees C dec 103.48 103.64 103.8 103.64 104.6 104.28 104.6 104.76 104.9 104.9	rees C degrees C	ees C deg 44.603 44.643 45.105 45.355 45.355 45.355 31.871 42.282 43.815 45.903 46.84 45.903 46.84 46.	rees C deg 41.619 41.663 41.766 41.776 41.775 41.793 42.292 47.058 50.947 52.842 62.168 78.357 90.481 95.004 97.689 98.745 99.371 99.994 99.371 99.994 99.371 00.26 100.46 100.26 100.46 100.65 100.84 101.04 101.23 101.62 101.62 101.89	rees C de 41.541 41.587 41.634 41.681 41.727 42.943 47.149 51.152 53.489 62.969 98.53 49.671 100.31 100.02 97.942 100.02 97.942 100.02 97.942 100.02 97.942 100.02 97.942	41.857 41.897 41.997 41.977 42.017 42.057 42.285 42.861 49.053 57.829 73.355 89.382 95.776 98.058 99.411 100.01 100.11 99.637 97.735 	42.859 42.899 42.94 42.98 43.02 43.06 43.323 43.894 50.57 59.538 74.991 91.101 97.397 99.511 100.6 101.25 101.72 101.25 98.961   T-150-10 degrees C 102.8 103.01 103.23 103.44 103.65 103.99 103.68 103.99 103.68 103.38	Grams/ml 1.1754 1.1754 1.1756 1.1746 1.1742 1.1737 1.1733 1.1729 1.1725 1.1711 1.1643 1.1607 1.1582 1.159 1.1633 1.1675 1.1715 1.1755 1.1794 1.1834  D-150-1 evaporator density Grams/ml 1.235 1.2469 1.2509 1.2549 1.2588 1.2628 1.2688 1.2628	137.02 136.99 136.97 136.94 136.92 136.89 136.87 136.84 136.82 137.05 137.39 136.35 132.79 128.99 125.09 120.8 116.68 113.27 123.08  L-150-1 evaporator level Inches 33.55 132.01 130.75 132.99 129.29 129.8 131.13	-2.732 -2.66 -2.587 -2.514 4.5011 434.46 -11.47 -10.86 1556.5 1722.1 1722.3 1733.4 1730.3 1734.7 1730.5 1731 1718 1713.3 1731  F350-1 evaporator steam flow b/hour 1738.9 1732.2 1727.5 1728.8 1726.9 1735.3 1724.9 1720.2 1729.2	74.848 1 74.825 74.802 74.78 74.757 74.735 74.712 74.689 74.71 74.765 74.822 74.875 74.929 74.984 75.039 75.094 75.149 75.204 75.259  T-336-1C degrees F 75.972 76.027 76.082 76.137 76.192 76.302 76.356 76.411	56.42 11 156.5 156.57 156.65 156.73 156.88 156.88 156.77 156.64 156.51 156.39 156.26 155.87 155.75 155.62 155.49 155.36	8.0286 8.0294 8.0309 8.0309 8.0317 8.0325 8.0333 11.451 11.445 11.442 11.436 11.433 11.431 11.425 11.422 11.419 11.37 11.38 11.37 11.37 11.37 11.37 11.37 11.37 11.37 11.37 11.37 11.37 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36 11.36	48.7 48.7	0.008 0.008	IN WC -0.056	IN WC 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	IN WC 2.0152 2.0151 2.0151 2.0155 2.015 2.015 2.0149 2.0149 2.0148 2.0147 2.0147 2.0146 2.0145 2.0145 2.0145 2.0145 2.0145 2.0145 2.0146 2.0136 2.0130 2.0139 2.0138 2.0137 2.0137 2.0136 2.0136 2.0136	IN WC 8.4546 8.4551 8.4557 8.4622 8.4667 8.4551 8.4551 8.4554 8.4551 8.4554 8.4405 8.4331 8.4294 8.4296 8.3773 8.3736 8.3692 8.3662 8.3662 8.3587 8.355 8.3513 8.3476	degrees F 183.33 183.33 183.33 183.32 183.32 183.32 183.32 183.32 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.31 183.37 183.27 183.27 183.27 183.27 183.27 183.27 183.27 183.27 183.27 183.27	degrees F 105.66 105.65 105.65 105.65 105.65 105.65 105.65 105.65 105.64 105.64 105.64 105.64 105.64 105.63 105.63 105.63 105.61 105.61 105.61 105.61 105.61	degrees F 105.56	98.865 98.866 98.867 98.869 98.87 98.871 98.873 98.874 98.875 98.876 98.878 98.881 98.883 98.884 98.885 98.884 98.885 98.888	degrees F sci 162.98 6: 162.96 6: 162.96 6: 162.96 6: 162.94 6: 162.94 6: 162.92 6: 162.92 6: 162.92 6: 162.92 6: 162.93 6: 162.89 6: 162.89 6: 162.89 6: 162.88 6: 162.88 6: 162.88 6: 162.88 6: 162.87 6: 162.77 6: 162.77 6: 162.77 6: 162.77 6: 162.75 6: 162.75 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.73 6: 162.71 6: 162.71 6:	ofm 635.69 614.51 649.32 601.31 652.47 634.15 627.71 661.54 661.68 654.57 661.54 661.5

		OR PARAMI																				_						
Time	T-150-1	T-150-2	T-150-3	T-150-4	T-150-5	T-150-6	T-150-7	T-150-8	T-150-9	T-150-10	D-150-1 evaporator density	L-150-1 evaporator level	F350-1 evaporator steam flow	T-336-1C	F 136-1	P-122-1	P-130-2 F	PD-130-1-1	PD-130-2-1	PD-130-3-1	PD-130-4-1 F	PD-130-1	T-335-2 T	-130-1-1 T	<b>r-130-2-1</b>	T-130-3-1 T	-130-4-1 F-130-1	
18 JUN 01 08:30:00	degrees C 59.957	degrees C 60.965	degrees C 66.25	degrees C d 66.442	legrees C o 47.646	degrees C o 46.878	degrees C 43,371	degrees C 43.595	degrees C 42.188	degrees C 44.149	Grams/ml 1.177	Inches 137.56	lb/hour -0.282	degrees F 75.329	scfm 151.75	"wcvac 8,8057	"wcvac 48,912	IN WC 0.0085	IN WC -0.06	IN WC	IN WC 2.053	IN WC 8.4825		degrees F 103.82	degrees F 102.96	degrees F 96.733	degrees F scfm 159.87 619.4	12
18 JUN 01 08:45:00	95.718	96.383	96.059	95.586	47.993	48.212	43.429	43.654	42.381	44.21	1.1764	137.61	4.8518	75.25	151.74	8.8073	48.913	0.0085	-0.059	-0.004 -0.003	2.053	8.4822	181.04	103.81	102.96	96.72	159.86 628.7	78
18 JUN 01 09:00:00 18 JUN 01 09:15:00	96.731 97.429	97.755 98.417	97.024 97.711	97.115 97.913	65.411 86.11	66.66 85.999	43.487 46.517	43.713 43.772	42.575 43.05	44.272 44.333	1.1758 1.1736	137.66 137.71	465.04 925.33	75.172 75.094	151.73 151.72	8.8088 8.8104	48.914 48.914	0.0085 0.0086	-0.059 -0.059	-0.003 -0.003	2.0531 2.0531	8.4818 8.4815	181.03 181.03	103.79 103.78	102.95 102.94	96.707 96.694	159.85 619.3 159.84 632.2	
18 JUN 01 09:30:00 18 JUN 01 09:45:00	98.127 98.678	99.08 99.398	98.399 99.041	98.71 99.055	94.438 95.942	93.726 95.51	53.309 64.029	43.831 47.432	47.983 58.866	49.769 60.832	1.1666 1.162	137.76 137.81	1283.3	75.016	151.7	8.8119	48.915	0.0086	-0.059	-0.003	2.0532	8.4811	181.02	103.77	102.93	96.68	159.83 602.9	91
18 JUN 01 10:00:00	99.167	99.714	99.369	99.377	95.478	95.098	80.904	53.887	75.85	77.923	1.1585	137.87	1742.2 1723.7	74.937 74.859	151.69 151.68	8.8134 8.815	48,915 48,916	0.0086 0.0086	-0.059 -0.059	-0.003 -0.003	2.0532 2.0533	8.4807 8.4804	181.02 181.02	103.75 103.74	102.93 102.92	96.667 96.654	159.82 616.2 159.81 596.4	19
18 JUN 01 10:15:00 18 JUN 01 10:30:00	99.657 100.15	100.02 100.24	99.697 100.14	99.738 100.47	94.277 93.246	94.312 94.083	91.523 96.219	64.716 81.45	91.039 96.627	91.368 98.162	1.1549 1.1586	135.54 131.74	1725.6 1727.7	74.781 74.702	151.67 151.65	8.8165 8.8181	48.917 48.917	0.0087 0.0087	-0.059 -0.059	-0.003 -0.003	2.0533 2.0534	8.48 8.4797	181.01 181.01	103.73 103.71	102.91 102.91	96.641 96.628	159.81 597.7 159.8 629.1	
18 JUN 01 10:45:00 18 JUN 01 11:00:00	100.64	100.45	100.81 101.49	101.2	94.625	94.675	98.347	92.713	98.872	100.08	1.1631	127.94	1731.2	74.624	151.64	8.8196	48.918	0.0087	-0.059	-0.003	2.0534	8.4793	181	103.7	102.9	96.615	159.79 651.0	)4
18 JUN 01 11:15:00	100.89 101.14	100.67 100.89	101.49	101.64 102	95.89 97.155	95.991 96.981	99.533 100.25	97.066 99.192	100.09 100.73	101.28 101.56	1.1674 1.1715	123.94 119.66	1728.8 1728	74.546 74.579	151.63 151.62	8.8212 8.8227	48.918 48.919	0.0087 0.0088	-0.059 -0.059	-0.003 -0.003	2.0535 2.0535	8.479 8.4786	181 180.99	103.69 103.68	102.89 102.88	96.602 96.589	159.78 630.8 159.77 624.9	
18 JUN 01 11:30:00	101.38	101.1	101.89	102.34	98.42	97.92	99.971	100.4	100.27	101.15	1.1757	115.59	1738.2	74.649	151.6	8.8242	48.92	0.0088	-0.059	-0.003	2.0536	8.4782	180.99	103.66	102.88	96.576	159.76 646.	.2
	EVAPORAT	OR PARAME	TERS, 0010	END-1							D-150-1	L-150-1	F350-1									_						
											evaporator	evaporator	evaporator															
Time					T-150-5 egrees C				T-150-9 degrees C	T-150-10 degrees C	density Grams/ml	level Inches	steam flow lb/hour	T-336-1C degrees F	F 136-1 P	<b>'-122-1</b> i "wcvac	P-130-2 P "wcvac	'D-130-1-1 F IN WC	PD-130-2-1 I IN WC	PD-130-3-1 I IN WC	PD-130-4-1 P IN WC	PD-130-1 IN WC	T-335-2 T degrees F	'-130-1-1 T degrees F	-130-2-1 1 degrees F	F-130-3-1 T degrees F	-130-4-1 F-130-1 degrees F scfm	
18 JUN 01 15:00:00 18 JUN 01 15:15:00	102.84 103.01	103.5 103.65	103.87 104.11	103.4 103.57	97.913 98.352	96.962 97.241	100.53 100.73	102.15 102.31	101.82 101.97	103.09 103.24	1.2337	132.18	1739.2	75.626	151.43	8.8459	48.928	0.0091	-0.059	-0.003	2.0542	8.4733	180.93	103.48	102.77	96.393	159.64 629.7	
18 JUN 01 15:30:00	103.19	103.79	104.26	103.74	98.638	97.519	100.92	102.46	102.12	103.39	1.2379 1.242	130.75 131.21	1731.1 1729.7	75.696 75.766	151.41 151.4	8.8474 8.849	48.929 48.929	0.0091 0.0092	-0.059 -0.059	-0.003 -0.003	2.0543 2.0543	8.4729 8.4725	180.92 180.92	103.47 103.45	102.77 102.76	96.38 96.367	159.63 625.4 159.62 630.6	
18 JUN 01 15:45:00 18 JUN 01 16:00:00	103.36 103.53	103.94 104.08	104.38 104.5	103.91 104.07	98.846 99.054	97.757 97.948	101.12 101.32	102.61 102.76	102.26 102.41	103.54 103.69	1.2462 1.2503	131.86 132.75	1731.7 1726.4	75.835 75.905	151.39 151.37	8.8505 8.852	48.93 48.93	0.0092 0.0092	-0.059 -0.059	-0.003 -0.003	2.0544 2.0544	8.4722 8.4718	180.92 180.91	103.44 103.43	102.75 102.75	96.354 96.341	159.61 614.2 159.6 629.5	
18 JUN 01 16:15:00	103.71	104.13	104.62	104.24	99.261	98.139	101.52	102.91	102.56	103.84	1.2545	132.59	1733.2	75.975	151.36	8.8536	48.931	0.0092	-0.059	-0.003	2.0545	8.4715	180.91	103.41	102.74	96.328	159.6 625.1	14
18 JUN 01 16:30:00 18 JUN 01 16:45:00	103.88 104.06	104.17 104.22	104.74 104.86	104.41 104.58	99.445 99.558	98.33 98.521	101.71 101.91	103.02 103.06	102.64 102.56	103.94 103.98	1.2586 1.2628	131.52 131.42	1735.1 1730.8	76.045 76.115	151.35 151.34	8.8551 8.8567	48.932 48.932	0.0093 0.0093	-0.059 -0.059	-0.003 -0.003	2.0545 2.0546	8.4711 8.4708	180.9 180.9	103.4 103.39	102.73 102.72	96.315 96.302	159.59 638.5 159.58 646.9	
18 JUN 01 17:00:00 18 JUN 01 17:15:00	104.23 104.1	104.26 104.3	104.98 105.06	104.75 104.74	99.671 99.783	98.712 98.903	102.11 102.31	103.11 103.15	102.48 102.39	104.03 104.07	1.2669 1.2711	131.32 131.22	1744 1728.4	76.184 76.254	151.32 151.31	8.8582 8.8598	48.933 48.933	0.0093 0.0093	-0.059 -0.059	-0.003 -0.003	2.0546 2.0547	8.4704 8.47	180.89 180.89	103.37 103.36	102.72 102.71	96.288 96.275	159.57 634.0 159.56 652.0	
18 JUN 01 17:30:00	103.93	104.35	104.86	104.56	99.896	99.094	102.51	103.2	102.31	104.12	1.2751	131.13	1729.2	76.324	151.3	8.8613	48.934	0.0094	-0.059	-0.003	2.0547	8.4697	180.89	103.35	102.7	96.262	159.55 642.6	<b>57</b>
18 JUN 01 17:45:00 18 JUN 01 18:00:00	103.76 103.58	104.39 104.44	104.66 104.46	104.38 104.19	100.01 100.12	99.285 99.476	102.49 102.33	103.24 103.28	102.22 102.14	104.16 104.21	1.2777 1.2803	131.03 130.93	1734.8 1736.5	76.394 76.464	151.29 151.27	8.8629 8.8644	48.934 48.935	0.0094 0.0094	-0.059 -0.059	-0.003 -0.003	2.0548 2.0548	8.4693 8.469	180.88 180.88	103.34 103.32	102.69 102.69	96.249 96.236	159.54 640.0 159.54 634.1	
	EVAPORAT	OR PARAME	TERS, 0010	STRT-2			<del> </del>				D-150-1	L-150-1	F350-1									_						
Time								T-150-8		T-150-10	evaporator density	evaporator level	evaporator steam flow	T-336-1C	F 136-1 P							PD-130-1	T-335-2 T				-130-4-1 F-130-1	
19 JUN 01 08:00:00	degrees C 57.26	degrees C 59.007	degrees C 66.803	degrees C de 6.205	egrees C o 46.013				-	T-150-10 degrees C 42.646	density	level	•	T-336-1C degrees F 76.791	F 136-1 P scfm 150.57	-122-1 F "wcvac 8.9509	P-130-2 P "wcvac 48.968	D-130-1-1 F IN WC 0.0107	PD-130-2-1 F IN WC -0.057	PD-130-3-1 I IN WC -0.001	PD-130-4-1 P IN WC 2.0575				-130-2-1 1 degrees F 102.28		<b>7-130-4-1 F-130-1</b> degrees F scfm 159.05 648.1	15
	degrees C	degrees C	degrees C	degrees C d	egrees C	degrees C d 44.578 49.344	egrees C	degrees C	degrees C 41.222 41.265	degrees C 42.646 42.685	density Grams/ml 1.1796 1.1792	level nches 136.63 136.59	steam flow b/hour 0.7685 5.2849	degrees F 76.791 76.797	scfm 150.57 150.55	"wcvac 8.9509 8.9524	*wcvac 48.968 48,969	IN WC 0.0107 0.0108	IN WC -0.057 -0.057	IN WC -0.001 -0.001	IN WC 2.0575 2.0575	IN WC 8.449 8.4487	degrees F 180.63 180.63	degrees F 102.59 102.58	degrees F 102.28 102.27	degrees F 95.505 95.492	degrees F scfm 159.05 648.1 159.04 646.7	'2
19 JUN 01 08:00:00 19 JUN 01 08:15:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00	degrees C 57.26 96.539 97.501 98.288	degrees C 59.007 97.633 98.482 99.086	degrees C 66.803 96.902 97.727 98.382	6.205 96.755 97.745 98.603	egrees C d 46.013 49.033 72.128 91.532	degrees C d 44.578 49.344 72.202 90.707	egrees C 41.262 41.303 42.046 46.395	degrees C 41.929 41.97 42.834 47.041	degrees C 41.222 41.265 41.307 42.405	degrees C 42.646 42.685 42.723 43.767	density  Grams/ml 1.1796 1.1792 1.1788 1.1761	level nches 136.63 136.59 136.54 136.5	steam flow b/hour 0.7685 5.2849 646.57 1020.2	degrees F 76.791 76.797 76.804 76.81	scfm 150.57 150.55 150.54 150.53	"wcvac 8.9509 8.9524 8.954 8.9555	"wcvac 48.968 48.969 48.97 48.97	IN WC 0.0107 0.0108 0.0108 0.0108	IN WC -0.057 -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576 2.0576	IN WC 8.449 8.4487 8.4483 8.4479	degrees F 180.63 180.63 180.62 180.62	degrees F 102.59 102.58 102.56 102.55	degrees F 102.28 102.27 102.27 102.26	degrees F 95.505 95.492 95.478 95.465	degrees F scfm 159.05 648.1 159.04 646.7 159.03 627.2 159.02 652.9	72 22 98
19 JUN 01 08:00:00 19 JUN 01 08:15:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00 19 JUN 01 09:00:00 19 JUN 01 09:15:00	degrees C 57.26 96.539 97.501 98.288 99.075 99.382	degrees C 59.007 97.633 98.482 99.086 99.69 100.02	degrees C 66.803 96.902 97.727 98.382 99.037 99.54	degrees C d 6.205 96.755 97.745 98.603 99.321 99.611	egrees C of 46.013 49.033 72.128 91.532 95.293 96.311	degrees C d 44.578 49.344 72.202 90.707 94.926 96.65	egrees C 41.262 41.303 42.046 46.395 54.169 66.713	degrees C 41.929 41.97 42.834 47.041 55.001 67.58	degrees C 41.222 41.265 41.307	degrees C 42.646 42.685 42.723	density Grams/ml 1.1796 1.1792 1.1788	level nches 136.63 136.59 136.54	steam flow b/hour 0.7685 5.2849 646.57	degrees F 76.791 76.797 76.804	scfm 150.57 150.55 150.54	"wcvac 8.9509 8.9524 8.954	*wcvac 48.968 48.969 48.97	IN WC 0.0107 0.0108 0.0108	IN WC -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576	IN WC 8.449 8.4487 8.4483	degrees F 180.63 180.63 180.62	degrees F 102.59 102.58 102.56	degrees F 102.28 102.27 102.27	degrees F 95.505 95.492 95.478	degrees F scfm 159.05 648.1 159.04 646.7 159.03 627.2	72 22 98 31
19 JUN 01 08:00:00 19 JUN 01 08:15:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00 19 JUN 01 09:00:00 19 JUN 01 09:15:00 19 JUN 01 09:30:00	degrees C 57.26 96.539 97.501 98.288 99.075 99.382 99.683	degrees C 59.007 97.633 98.482 99.086 99.69 100.02 100.32	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833	6.205 96.755 97.745 98.603 99.321 99.611 99.901	egrees C 0 46.013 49.033 72.128 91.532 95.293 96.311 96.681	degrees C d 44.578 49.344 72.202 90.707 94.926 96.65 96.119	egrees C 41.262 41.303 42.046 46.395 54.169 66.713 84.552	degrees C 41.929 41.97 42.834 47.041 55.001 67.58 84.578	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403	degrees C 42.646 42.685 42.723 43.767 49.553 62.257 80.84	density	level nches 136.63 136.59 136.54 136.5 136.55 136.82 136.82	steam flow  b/hour	degrees F 76.791 76.797 76.804 76.816 76.822 76.829	scfm 150.57 150.55 150.54 150.53 150.52 150.5 150.49	"wcvac 8.9509 8.9524 8.954 8.9555 8.9571 8.9586 8.9602	*wevac 48.968 48.969 48.97 48.97 48.971 48.971 48.972	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109	IN WC -0.057 -0.057 -0.057 -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0577 2.0577 2.0577	IN WC 8.449 8.4487 8.4483 8.4479 8.4476 8.4472 8.4469	degrees F 180.63 180.63 180.62 180.62 180.61 180.61	degrees F 102.59 102.58 102.56 102.55 102.54 102.53 102.51	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.24	degrees F 95.505 95.492 95.478 95.465 95.452 95.439 95.426	degrees F         scfm           159.05         648.1           159.04         646.7           159.03         627.2           159.02         652.9           159.01         628.3           158.99         634.9	72 22 98 31 35
19 JUN 01 08:00:00 19 JUN 01 08:15:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00 19 JUN 01 09:00:00 19 JUN 01 09:30:00 19 JUN 01 09:30:00 19 JUN 01 09:45:00 19 JUN 01 10:00:00	degrees C 57.26 96.539 97.501 98.288 99.075 99.382 99.683 99.984 100.22	degrees C 59.007 97.633 98.482 99.086 99.69 100.02 100.32 100.63 100.88	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13 100.4	degrees C d 6.205 96.755 97.745 98.603 99.321 99.611 99.901 100.19	egrees C c 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.492 93.555	degrees C d 44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511 93.411	egrees C 41.262 41.303 42.046 46.395 54.169 66.713 84.552 92.692 96.919	degrees C 41.929 41.97 42.834 47.041 55.001 67.58 84.578 94.104 97.69	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403 92.171 97.199	degrees C 42.646 42.685 42.723 43.767 49.553 62.257 80.84 94.175 98.671	density Grams/ml 1.1796 1.1792 1.1788 1.1761 1.177 1.1667 1.1639 1.1611 1.1643	level nches 136.63 136.59 136.54 136.55 136.55 136.82 136.82 136.84 134.53 131.01	steam flow b/hour 0.7685 5.2849 646.57 1020.2 1429.7 1727.2 1739.5 1734.5 1737.2	degrees F 76.791 76.797 76.804 76.81 76.816 76.822 76.829 76.835 76.841	scfm 150.57 150.55 150.54 150.53 150.52 150.5 150.49 150.48 150.46	"wcvac 8.9509 8.9524 8.954 8.9555 8.9571 8.9586 8.9602 8.9617 8.9632	*wevac 48.968 48.969 48.97 48.97 48.971 48.971 48.972 48.972 48.973	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109 0.0109 0.0109	IN WC -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0577 2.0577 2.0577 2.0577 2.0578	IN WC 8.449 8.4487 8.4483 8.4479 8.4476 8.4472 8.4469 8.4465 8.4462	degrees F 180.63 180.63 180.62 180.62 180.61 180.61 180.61 180.6 180.6	degrees F 102.59 102.58 102.56 102.55 102.54 102.53 102.51 102.5 102.49	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.24 102.23 102.22	degrees F 95.505 95.492 95.478 95.465 95.452 95.439 95.426 95.413 95.4	degrees F         scfm           159.05         648.1           159.04         646.7           159.03         627.2           159.01         682.9           159.01         628.3           158.99         634.9           158.99         657.2           158.98         665.2           688.99         665.2	72 22 98 31 35 97 28
19 JUN 01 08:00:00 19 JUN 01 08:15:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00 19 JUN 01 09:00:00 19 JUN 01 09:15:00 19 JUN 01 09:30:00 19 JUN 01 09:45:00	degrees C 57.26 96.539 97.501 98.288 99.075 99.382 99.683 99.984	degrees C 59.007 97.633 98.482 99.086 99.69 100.02 100.32 100.63	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13	degrees C d 6.205 96.755 97.745 98.603 99.321 99.611 99.901 100.19	egrees C d 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.492	44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511	egrees C 41.262 41.303 42.046 46.395 54.169 66.713 84.552 92.692	41.929 41.97 42.834 47.041 55.001 67.58 84.578 94.104	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403 92.171	degrees C 42.646 42.685 42.723 43.767 49.553 62.257 80.84 94.175	density Grams/ml 1.1796 1.1792 1.1788 1.1761 1.17 1.1667 1.1639 1.1611	level nches 136.63 136.59 136.54 136.55 136.55 136.82 136.82 136.84 134.53	steam flow b/hour 0.7685 5.2849 646.57 1020.2 1429.7 1727.2 1739.5 1734.5	degrees F 76.791 76.797 76.804 76.81 76.816 76.822 76.829 76.835	scfm 150.57 150.55 150.54 150.53 150.52 150.5 150.49 150.48	"wcvac 8.9509 8.9524 8.954 8.9555 8.9571 8.9586 8.9602 8.9617	*wevac 48.968 48.969 48.97 48.97 48.971 48.971 48.972 48.972	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109	IN WC -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0577 2.0577 2.0577 2.0578	IN WC 8.449 8.4487 8.4483 8.4479 8.4476 8.4472 8.4469 8.4465	degrees F 180.63 180.63 180.62 180.62 180.61 180.61 180.61	degrees F 102.59 102.58 102.56 102.55 102.54 102.53 102.51 102.5	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.24 102.23	degrees F 95.505 95.492 95.478 95.465 95.452 95.439 95.426 95.413	degrees F         scfm           159.05         648.1           159.04         646.7           159.03         627.2           159.01         628.3           159         635.8           158.99         634.9           158.99         657.2	72 22 98 91 95 97 98 97
19 JUN 01 08:00:00 19 JUN 01 08:30:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00 19 JUN 01 09:00:00 19 JUN 01 09:30:00 19 JUN 01 09:30:00 19 JUN 01 10:00:00 19 JUN 01 10:00:00 19 JUN 01 10:15:00 19 JUN 01 10:15:00 19 JUN 01 10:45:00 19 JUN 01 10:45:00	degrees C 57.26 96.539 97.501 98.288 99.075 99.382 99.683 99.984 100.22 100.45 100.68 100.91	degrees C 59.007 97.633 98.482 99.086 99.69 100.02 100.32 100.63 100.88 101.1 101.32 101.54	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13 100.63 100.63 100.86	degrees C d 6.205 96.755 97.745 98.603 99.321 99.611 99.901 100.19 100.41 100.64 100.87	egrees C d 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.492 93.555 94.701 95.75 96.799	degrees C d 44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511 93.411 94.615 95.721 96.827	egrees C 41.262 41.303 42.046 46.395 54.169 66.713 84.552 92.692 96.919 98.574 99.633 100.29	degrees C 41,929 41,97 42,834 47,041 55,001 67,58 84,578 94,104 97,69 99,408 100,56 101,06	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403 92.171 97.199 99.048 100.17 100.52	degrees C 42.646 42.685 42.763 43.767 49.553 62.257 80.84 94.175 98.671 100.38 101.55 102.08	density Grams/ml 1.1796 1.1792 1.1788 1.1761 1.176 1.1667 1.1639 1.1611 1.1643 1.1684 1.1733 1.1789	Ievel nches 136.63 136.59 136.54 136.55 136.55 136.82 136.84 134.53 131.01 127.04 123.13 119.42	steam flow b/hour 0.7685 5.2849 646.57 1020.2 1429.7 1727.2 1739.5 1734.5 1737.2 1724.6 1733 1745.2	degrees F 76.791 76.804 76.814 76.816 76.822 76.829 76.835 76.841 76.847 76.853	scfm 150.57 150.55 150.54 150.53 150.52 150.49 150.48 150.46 150.45 150.44 150.43	"wcvac 8.9509 8.9524 8.954 8.9555 8.9571 8.9586 8.9602 8.9617 8.9632 8.9648 8.9663 8.9679	"wevac 48.968 48.969 48.97 48.97 48.971 48.972 48.972 48.973 48.974 48.974 48.975	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109 0.0109 0.0109 0.0109 0.01109 0.01109 0.0111	IN WC -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0577 2.0577 2.0577 2.0577 2.0578 2.0578 2.0579 2.0579	8.449 8.4487 8.4483 8.4479 8.4476 8.4472 8.4469 8.4465 8.4462 8.4455 8.4455	degrees F 180.63 180.63 180.62 180.62 180.61 180.61 180.6 180.5 180.59 180.59	degrees F 102.59 102.58 102.56 102.55 102.54 102.53 102.51 102.5 102.49 102.47 102.46	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.23 102.22 102.22 102.21	degrees F 95.505 95.492 95.478 95.465 95.452 95.439 95.426 95.413 95.4 95.387 95.374	degrees F         scfm           159.05         648.1           159.04         648.7           159.03         627.2           159.01         682.9           159.01         628.3           158.99         634.9           158.99         657.2           158.97         658.1           158.96         626.9           158.97         658.1           158.95         655.6	72 22 28 31 35 37 28 27 39 44
19 JUN 01 08:00:00 19 JUN 01 08:15:00 19 JUN 01 08:30:00 19 JUN 01 08:45:00 19 JUN 01 09:00:00 19 JUN 01 09:30:00 19 JUN 01 09:45:00 19 JUN 01 10:45:00 19 JUN 01 10:15:00 19 JUN 01 10:30:00 19 JUN 01 10:30:00 19 JUN 01 11:30:00 19 JUN 01 11:15:00 19 JUN 01 11:15:00	degrees C 57.26 96.539 97.501 98.288 99.075 99.382 99.683 99.984 100.22 100.45 100.68 100.91 101.14	degrees C 59.007 97.633 98.482 99.086 99.69 100.02 100.63 100.88 101.1 101.32 101.54 101.76	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13 100.63 100.63 101.08 101.08 101.31	6.205 96.755 97.745 98.603 99.321 99.901 100.19 100.41 100.64 100.87 101.09 101.32 101.55	egrees C d 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.492 93.555 94.701 95.75 96.799 97.847 96.682	44.578 44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511 93.411 94.615 95.721 96.827 97.933 96.688	egrees C 41.262 41.303 42.046 46.395 54.169 66.713 84.552 92.692 96.919 98.574 99.633 100.29 100.04 97.87	degrees C 41,929 41,97 42,834 47,041 55,001 67,58 84,578 94,104 97,69 99,408 100,56 101,06 101,5 98,598	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403 92.171 97.199 99.048 100.17 100.52 100.13 98.398	degrees C 42.646 42.685 42.723 43.767 49.553 62.257 80.84 94.175 98.671 100.38 101.55 102.08 102.54 99.664	density  Grams/ml  1.1796  1.1792  1.1788  1.1761  1.17  1.1667  1.1633  1.1643  1.1733  1.1789  1.1845	level nches 136.63 136.59 136.54 136.55 136.55 136.84 134.53 131.01 127.04 123.13 119.42 115.71 121.08	steam flow b/hour 0.7685 5.2849 646.57 1020.2 1429.7 1727.2 1739.5 1734.5 1737.2 1724.6 1733 1745.2 1733.5	degrees F 76.791 76.804 76.816 76.822 76.829 76.835 76.847 76.853 76.866 76.866	scfm 150.57 150.55 150.54 150.53 150.52 150.5 150.49 150.48 150.45 150.44 150.43	"wcvac 8.9509 8.9524 8.9554 8.9555 8.9571 8.9586 8.9602 8.9617 8.9632 8.9648 8.9663 8.9679 8.9694 8.971	"wevac 48.968 48.969 48.97 48.97 48.971 48.972 48.972 48.973 48.974 48.974 48.975 48.975	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109 0.0109 0.0109 0.0109 0.0109	IN WC -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057 -0.057	IN WC -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001 -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0577 2.0577 2.0577 2.0578 2.0578 2.0579	8.449 8.4487 8.4483 8.4479 8.4476 8.4472 8.4469 8.4465 8.4465 8.4458	degrees F 180.63 180.62 180.62 180.61 180.61 180.6 180.6 180.5 180.59 180.59 180.58	degrees F 102.59 102.58 102.56 102.55 102.51 102.51 102.5 102.47 102.46 102.45 102.43	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.23 102.22 102.22 102.21 102.2 102.2 102.1 102.19	degrees F 95.505 95.492 95.478 95.465 95.452 95.439 95.426 95.413 95.347 95.374 95.361 95.348 95.335	degrees F         scfm           159.05         648.1           159.04         646.7           159.03         627.2           159.02         652.9           159.01         628.3           158.99         634.9           158.99         657.2           158.98         665.2           158.97         658.1           158.95         655.6           158.96         626.9           158.97         658.1           158.96         626.9           158.97         658.4           688.94         620.6           158.93         608.9	72 22 28 31 35 37 28 8 27 9 9 9 9 9 9 9
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104.7 104.5 104.5 104.5 104.5 104.5 104.5 104.5 105.5 105.2 105.5	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13 100.63 100.86 101.08 101.31 101.54 101.77 101.99 CTERS, 00101 T-150-3 degrees C 103.94 104.12 104.3 104.48 105.02 105.37	Gegrees C de 6.205 96.755 97.745 98.603 99.321 99.611 99.901 100.19 100.41 100.64 100.87 101.09 101.32 101.55 101.78 102 END-2  T-150-4 Jegrees C de 103.92 104.45 104.45 104.45 104.87 105.15 105.32	egrees C d 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.492 93.555 94.701 95.75 96.799 97.847 96.682 97.159  7-150-5 egrees C d 97.639 98.114 98.588 99.025 99.305 99.707 100.035 100.39 100.73	44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511 94.615 95.721 96.8827 97.933 96.688 96.042 97.065  7-150-6 egrees C d 97.229 97.692 98.155 98.618 98.966 99.3 99.634 99.968 100.3	egrees C 41.262 41.303 42.046 46.395 54.169 96.919 98.574 99.639 100.04 97.87 99.056 99.7	degrees C 41,929 41,97 42,834 47,041 55,001 67,58 84,578 94,104 97,69 99,408 100,56 101,06 101,5 98,598 99,824 100,69 T-150-8 degrees C 101,58 102,63 102,77 102,91 103,06 103,2 103,34 103,49 103,76	degrees C 41,222 41,265 41,307 42,405 48,917 61,709 79,403 92,171 97,199 99,048 100,17 100,52 100,13 98,398 99,682 100,45 T-150-9 degrees C 101,237 102,5 102,64 102,77 102,9 103,04 103,17 103,36	degrees C 42.646 42.646 42.685 42.723 43.767 49.553 62.257 80.84 94.175 98.671 100.38 101.55 102.08 102.54 99.664 100.7 101.57  T-150-10 degrees C 103.08 103.56 103.7 103.85 104 104.15 104.29 104.46 104.74	density   Grams/m    1.1796   1.1792   1.1788   1.1761   1.17   1.1667   1.1639   1.1611   1.1643   1.1643   1.1733   1.1789   1.1845   1.1901   1.1956   1.2012     D-150-1   evaporator density   Grams/m    1.2347   1.2347   1.2347   1.2426   1.2465   1.2504   1.25543   1.2575   1.2607   1.2639	Ievel	steam flow  b/hour   0.7685   5.2849   646.57   1020.2   1429.7   1727.2   1739.5   1734.5   1735.2   1736.2   1739.2   1739.5   1730.2   1729.3     F350-1 evaporator steam flow  b/hour   1731.5   1729.8   1729.2   1739.4   1729.8   1729.5   1729.5   1730.8   1729.5   1730.8   1729.5   1730.8   1730	76.96 76.97 76.804 76.814 76.815 76.822 76.829 76.829 76.835 76.841 76.847 76.853 76.866 76.872 76.878 76.884 76.894 76.947 76.953 76.965 76.965 76.971 76.971	scfm 150.57 150.55 150.54 150.53 150.52 150.48 150.44 150.43 150.41 150.43 150.39 150.38 150.26 150.26 150.22 150.22 150.22 150.21 150.2 150.19 150.16	"wcvac 8,9509 8,9524 8,954 8,9555 8,9571 8,9586 8,9602 8,9613 8,9648 8,9663 8,971 8,9724 8,988 8,989 8,991 8,9924 8,9957 8,9972 8,9984 8,9957 8,9972 8,9982 8,9982 8,99003	**Wevac	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109 0.0109 0.0109 0.0111	IN WC -0.057	IN WC -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0576 2.0577 2.0577 2.0577 2.0577 2.0578 2.0578 2.0578 2.0579 2.0581 2.0581 2.0581 2.0582  PD-130-4-1 IN WC 2.0586 2.0587 2.0587 2.0587 2.0588 2.0588 2.0589 2.0588	IN WC 8.449 8.4487 8.4487 8.4472 8.4469 8.4472 8.4469 8.4465 8.4465 8.4451 8.4447 8.4444 8.4437  IN WC 8.4401 8.4398 8.4398 8.4398 8.4387 8.4388 8.4376	degrees F 180.63 180.63 180.62 180.62 180.61 180.61 180.61 180.61 180.59 180.59 180.58 180.58 180.58 180.58 180.57  T-335-2 degrees F 180.53 180.57  T-385-2 180.51 180.51 180.51 180.51 180.51 180.55	degrees F 102.59 102.56 102.55 102.55 102.53 102.51 102.57 102.49 102.47 102.46 102.45 102.43 102.42 102.41 102.39  -130-1-1	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.24 102.22 102.22 102.21 102.19 102.19 102.17  -130-2-1 degrees F 102.11 102.1 102.0 102.08 102.08 102.06 102.06	degrees F 95.505 95.492 95.478 95.465 95.426 95.426 95.439 95.426 95.337 95.361 95.361 95.322 95.309 F-130-3-1 T degrees F 95.191 95.178 95.152 95.139 95.126 95.139 95.126 95.113 95.113 95.113 95.126 95.139	degrees F         scfm           159.04         648.1           159.03         648.1           159.03         627.2           159.01         628.3           159.05         655.8           158.99         634.9           158.99         657.2           158.97         658.1           158.98         665.2           158.95         655.6           158.93         608.9           158.93         648.4           158.93         643.8           458.93         643.8           158.80         665.2           158.81         665.2           158.82         630.8           158.83         647.2           158.84         665.8           158.85         665.2           158.79         666.2           158.79         666.2           158.79         665.2           158.78         652.6           158.77         650.3	72 228 311 35 77 88 77 9 34 34 32 93 11 36 6 6 11 22 26 6
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106.11	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13 100.63 100.86 101.08 101.31 101.54 101.77 101.99 TERS, 00101 T-150-3 degrees C 103.94 104.12 104.3 104.48 105.02 105.37 105.55 105.73 105.91	Gegrees C de 6.205 96.755 97.745 98.603 99.321 99.611 99.901 100.41 100.64 100.87 101.09 101.32 101.55 101.78 102 END-2  END-2  T-150-4 104.27 104.45 104.45 104.45 104.87 105.5 105.65 105.65 105.65 105.65	egrees C d 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.492 93.555 94.701 95.75 96.799 97.847 96.682 97.639 98.114 98.588 99.025 99.365 99.707 100.035 100.73 101.08 101.42 101.41	degrees C d 44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511 94.615 95.721 96.827 97.933 96.688 96.042 97.065  T-150-6 egrees C d 97.229 98.155 98.618 98.966 99.3 99.634 99.968 100.3 100.64 100.97 101.22	egrees C 41.262 41.303 42.046 46.395 54.169 96.919 98.574 99.639 100.04 97.87 99.056 99.7 T-150-7 egrees C 6 100.98 102.41 102.54 102.79 102.79 102.79 103.31 103.34 103.35	41,929 41,929 41,97 42,834 47,041 55,001 67,58 84,578 94,104 97,69 99,408 100,56 101,06 101,5 98,598 99,824 100,69 T-150-8 iegrees C 101,58 102,77 102,91 103,06 103,2 103,34 103,76 104,04 104,04 104,03 104,62	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403 92.171 97.199 99.048 100.17 100.52 100.13 98.398 99.682 100.45	degrees C 42.646 42.646 42.685 42.723 43.767 49.553 62.257 80.84 94.175 98.671 100.38 101.55 102.08 102.54 99.664 100.7 101.57  T-150-10 degrees C 103.08 103.56 103.7 103.85 104 104.15 104.29 105.31 105.59	Company	Ievel	steam flow b/hour 0.7685 5.2849 646.57 1020.2 1429.7 1727.2 1739.5 1734.5 1733.2 1724.6 1733.5 1723.2 1730.2 1729.3  F350-1 evaporator steam flow b/hour 1731.5 1729.8 1729.8 1729.8 1729.5 1730.8 1729.5 1730.8 1729.4	76.94 76.95 76.804 76.814 76.816 76.822 76.829 76.835 76.841 76.853 76.866 76.878 76.868 76.878 76.884 76.894 76.953 76.953 76.953 76.953 76.965 76.971 76.971 76.984 76.994 76.996 77.003	scfm 150.57 150.55 150.54 150.53 150.52 150.48 150.44 150.43 150.41 150.43 150.39 150.38 F 136-1 P scfm 150.26 150.22 150.21 150.22 150.21 150.2 150.19 150.15 150.16 150.15 150.14 150.12	"wcvac 8,9509 8,9524 8,954 8,9555 8,9571 8,9586 8,9602 8,9617 8,9632 8,9648 8,9674 8,9725 8,9741 F"wcvac 8,988 8,9895 8,991 8,9924 8,9957 8,9972 8,9972 8,9972 8,9972 8,9972 8,9903 9,0019 9,0034 9,0049	**Wevac 48.968 48.969 48.97 48.97 48.971 48.971 48.972 48.973 48.973 48.974 48.975 48.975 48.975 48.977 48.977  **Wevac 48.983 48.983 48.983 48.984 48.985 48.986 48.987 48.987 48.987 48.987 48.988	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109 0.0109 0.0109 0.01115 0.0115 0.01115 0.01115 0.01115	IN WC -0.057 -0.056	IN WC -0.001	IN WC 2.0575 2.0576 2.0576 2.0576 2.0577 2.0577 2.0577 2.0577 2.0578 2.0578 2.0578 2.0579 2.0581 2.0581 2.0581 2.0582  PD-130-4-1 IN WC 2.0586 2.0587 2.0587 2.0588 2.0588 2.0588 2.0588 2.0589 2.0589 2.0589 2.0589 2.0589 2.0589	IN WC 8.449 8.449 8.4487 8.4483 8.4472 8.4462 8.4462 8.4465 8.4465 8.4451 8.4447 8.4444 8.4437  IN WC 8.4401 8.4398 8.4394 8.4394 8.4398 8.4398 8.4398 8.4376 8.4373 8.4369 8.4369	degrees F 180.63 180.62 180.62 180.62 180.61 180.61 180.61 180.61 180.59 180.59 180.58 180.58 180.58 180.57  T-335-2 degrees F 180.53 180.52 180.51 180.51 180.51 180.51 180.51 180.51 180.51 180.51 180.54 180.49 180.49 180.48	degrees F 102.59 102.58 102.56 102.55 102.54 102.53 102.51 102.49 102.47 102.46 102.45 102.43 102.42 102.41 102.39  -130-1-1 T degrees F 102.28 102.26 102.25 102.21 102.21 102.21 102.15 102.15 102.15	degrees F 102.28 102.27 102.27 102.26 102.25 102.24 102.24 102.22 102.21 102.22 102.17 102.19 102.19 102.17  -130-2-1 102.17  -130-2-1 102.09 102.08 102.07 102.06 102.05 102.03 102.03	degrees F 95.505 95.492 95.478 95.465 95.429 95.426 95.439 95.413 95.37 95.361 95.335 95.322 95.309 F-130-3-1 T degrees F 95.191 95.178 95.152 95.139 95.126 95.139 95.126 95.139 95.139 95.139 95.139 95.139	degrees F   548.1   159.05   648.1   159.04   648.7   159.03   627.2   159.01   628.3   159.01   628.3   159.01   628.3   158.99   634.9   655.2   158.97   658.1   158.95   655.6   158.94   620.6   158.93   648.8   158.93   648.8   158.93   643.8   158.82   630.8   158.83   647.2   158.84   662.8   158.81   662.8   158.87   662.8   158.79   666.2   158.79   666.2   158.79   666.2   158.79   666.2   158.79   666.2   158.79   665.3   158.76   650.3   158.76   650.3   158.76   655.3   655.3   655.3   655.3	72 22 28 11 15 77 28 77 9 14 14 12 19 11 19 3 3 3 11 16 6 6 11 22 26 6 5 5 5 11
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105.58 105.58 105.58 105.76 105.93	degrees C 66.803 96.902 97.727 98.382 99.037 99.54 99.833 100.13 100.63 101.08 101.31 101.54 101.77 101.99 CTERS, 00101 T-150-3 degrees C 103.94 104.12 104.3 104.48 104.66 104.84 105.02 105.19 105.37 105.55 105.73	Gegrees C de 6.205 96.755 97.745 98.603 99.321 99.611 99.901 100.41 100.64 100.87 101.09 101.32 101.55 101.78 102 END-2  END-2  T-150-4  Gegrees C de 103.92 104.1 104.27 104.45 104.62 104.8 104.97 105.15 105.52 105.5 105.57 105.5	egrees C d 46.013 49.033 72.128 91.532 95.293 96.311 96.681 94.701 95.75 94.701 95.75 95.79 97.847 96.549 97.159  F-150-5 egrees C d 97.639 98.114 98.588 99.707 100.05 100.39 100.73 101.08 101.42	degrees C d 44.578 49.344 72.202 90.707 94.926 96.65 96.119 94.511 93.411 94.615 95.721 96.827 97.933 96.688 96.042 97.065  T-150-6 egrees C d 97.229 98.155 98.618 98.966 99.3 99.634 99.968 100.3 100.64 100.97	41.262 41.262 41.303 42.046 46.395 54.169 96.713 84.552 96.919 98.574 99.652 99.056 99.7 7 100.04 97.87 99.056 99.7 7 102.94 102.41 102.67 102.92 103.05 103.17 103.33 103.34	degrees C 41,929 41,97 42,834 47,041 55,001 67,58 84,578 94,104 97,69 99,408 100,56 101,06 101,5 98,598 4100,69 T-150-8 Jegrees C 101,58 102,63 102,77 102,91 103,06 103,24 103,34 103,49 103,76 104,04 104,33	degrees C 41.222 41.265 41.307 42.405 48.917 61.709 79.403 92.171 97.199 99.048 100.17 100.52 100.13 98.398 99.682 100.45  T-150-9 degrees C 101.32 102.37 102.5 102.64 102.77 102.9 103.04 103.17 103.36 103.64 103.92	degrees C 42.646 42.685 42.623 43.767 49.553 62.257 80.84 94.175 98.671 100.38 101.55 102.08 102.54 99.664 100.7 101.57  T-150-10  degrees C 103.08 103.56 103.7 103.85 104 104.15 104.46 104.74 105.02 105.31	1.796   1.1796   1.1796   1.1792   1.1788   1.1761   1.177   1.1667   1.1639   1.1611   1.1643   1.1643   1.1684   1.1733   1.1789   1.1845   1.1901   1.1956   1.2012     1.2	Ievel	steam flow b/hour 0.7685 5.2849 646.57 1020.2 1429.7 1727.2 1739.5 1734.5 1737.2 1724.6 1733.3 1745.2 1730.2 1729.3  F350-1 evaporator steam flow b/hour 1731.5 1729.8 1716.5 1729.2 1730.8 1729.6 1729.5 1730.8 1729.8 1747.8	76.91 76.797 76.804 76.814 76.816 76.822 76.829 76.835 76.841 76.847 76.866 76.872 76.878 76.884 76.984 76.959 76.959 76.978 76.978 76.984 76.984 76.998	scfm 150.57 150.55 150.54 150.53 150.52 150.48 150.44 150.43 150.44 150.39 150.38 F 136-1 P scfm 150.25 150.24 150.25 150.24 150.25 150.24 150.25 150.24 150.25 150.27 150.26 150.27 150.26 150.27 150.16 150.17 150.16 150.15 150.14	"wcvac 8,9509 8,9524 8,9554 8,9555 8,9571 8,9602 8,9617 8,9632 8,9648 8,9669 8,9725 8,9741 F"wcvac 8,988 8,9895 8,991 8,9926 8,991 8,9927 8,9928 9,0003 9,0019 9,0034	**Wevac	IN WC 0.0107 0.0108 0.0108 0.0108 0.0108 0.0109 0.0109 0.0109 0.0109 0.0111	IN WC -0.057	IN WC -0.001	IN WC 2.0575 2.0575 2.0576 2.0576 2.0576 2.0577 2.0577 2.0577 2.0577 2.0578 2.0578 2.0579 2.0579 2.058 2.0581 2.0581 2.0582  PD-130-4-1 IN WC 2.0586 2.0587 2.0587 2.0587 2.0588 2.0588 2.0588 2.0588 2.0588 2.0588 2.0588 2.0588 2.0588 2.0589 2.0589 2.0599 2.0599 2.0599	IN WC 8.449 8.4487 8.4487 8.4483 8.4476 8.4472 8.4469 8.4465 8.4465 8.4451 8.4447 8.4444 8.4437  PD-130-1 IN WC 8.4405 8.4405 8.4498 8.4394 8.4398 8.4398 8.4398 8.4398 8.4393 8.4383 8.4373 8.4369	degrees F 180.63 180.63 180.62 180.62 180.61 180.61 180.61 180.61 180.59 180.59 180.58 180.58 180.58 180.57  T-335-2 T degrees F 180.57  T-380.57  T-380.51 180.51 180.51 180.51 180.51 180.51 180.5 180.49 180.49 180.49	degrees F 102.59 102.58 102.56 102.55 102.53 102.51 102.51 102.51 102.49 102.47 102.42 102.43 102.42 102.41 102.39 Talegrees F 102.28 102.26 102.25 102.24 102.25 102.21 102.21 102.21	degrees F 102.28 102.27 102.27 102.26 102.24 102.24 102.23 102.22 102.22 102.21 102.19 102.19 102.17  -130-2-1	degrees F 95.505 95.492 95.478 95.465 95.452 95.439 95.443 95.341 95.361 95.361 95.322 95.309 F-130-3-1 T degrees F 95.178 95.182 95.182 95.191 95.178 95.152 95.139 95.152 95.139 95.152 95.139 95.152 95.139 95.152 95.139 95.152 95.139	degrees F   56m   159.05   648.1   159.04   648.7   159.02   652.9   159.02   652.9   655.2   158.99   657.2   158.95   655.6   158.94   620.6   158.93   648.4   158.93   648.4   158.93   648.8   158.83   647.2   649.7   158.83   647.2   158.84   655.6   158.84   665.8   158.85   665.8   158.87   666.5   158.79   666.5   158.79   666.5   158.79   666.5   158.79   666.5   158.79   666.5   158.79   665.6   158.77   650.3   158.76   652.6   158.77   650.3   158.76   665.6   158.77   650.3   158.76   665.6   158.77   650.3   158.76   650.3   158.	72 22 8 1 1 5 7 7 8 7 7 9 4 4 4 2 9 1 1 9 7 3 3 1 1 6 6 6 1 2 2 6 6 5 5 5 1 1 8 8

Table D-1. Evaporato	er parameters. EVAPORAT	OR PARAME	TERS, 0031	ISTRT-1																							
											D-150-1 evaporator	L-150-1 evaporator	F350-1 evaporator														
Time	T-150-1 degrees C					T-150-6 degrees C	T-150-7	T-150-8 degrees C	T-150-9 degrees C	T-150-10 degrees C	density	level Inches	steam flow	T-336-1C	F 136-1			PD-130-1-1 IN WC	PD-130-2-1 IN WC	PD-130-3-1	PD-130-4-1 IN WC	PD-130-1 IN WC	T-335-2 degrees F	T-130-1-1 degrees F	T-130-2-1 degrees F	<b>T-130-3-1</b> degrees F	T-130-4-1 F-130-1 degrees F scfm
20 JUN 01 08:00:00	54.996	56,163	64.408	64.363	40.55	39.642	32.805	33.555	34.667	36.002	1.1964	136.89	-7.56	degrees F 74.085	152.45	"wcvac 8.9506	"wcvac 48.8	0.008	-0.056		0 2.115	9 8.6697	180.67	102.14	101.	9 95.541	158.83 623.9
20 JUN 01 08:15:00 20 JUN 01 08:30:00	94.28 98.273	94.079 99.275	95.593 98.793	94.447 98.658	41.155 54.666	40.27 53.947	33.189 33.574	33.939 34.324	34.902 35.136	36.238 36.474	1.1932 1.19	136.77 136.65	-0.378 126.83	74.133 74.18	152.81 153.17	9.0915 9.2324	48.799 48.798	800.0 800.0	-0.056 <b>-</b> 0.056		0 2.115			102.15 102.16			
20 JUN 01 08:45:00 20 JUN 01 09:00:00	98.844 99.248	99.847 100.12	99.32 99.646	99.195 99.537	83.446 94.975	84.276 94.458	36.241 42.124	36.926 42.857	35.371 37.759	36.711 39.66	1.1858 1.1781	136.53 136.41	834.14 1284.2	74.227 74.275	153.53	9.3733 9.5141	48.798 48.797	0.008	-0.056	<b>;</b>	0 2.115 0 2.115			102.17 102.17			
20 JUN 01 09:15:00	99.651	100.39	99.971	99.878	97.217	96.681	51.938	52.692	46.365	48.09	1.1706	136.89	1711.7	74.322	153.89 154.25	9.655	48.796	0.008 0.008	-0.056 -0.056		0 2.115	2 8.665	180.67	102.18	3 101.9	5 95.549	158.84 645.89
20 JUN 01 09:30:00 20 JUN 01 09:45:00		100.66 100.9	100.3 100.58	100.22 100.49	97.347 96.936	97.698 97.164	67.03 86.333	67.738 86.044	60.963 81.417	62.193 83.347	1.1664 1.1622	137.32 137.28	1726.9 1743	74.369 74.417	154.61 154.97	9.5319 9.3324	48.796 48.795	0.008 0.008	-0.056 -0.056		0 2.11 0 2.114			102.19 102.2			
20 JUN 01 10:00:00 20 JUN 01 10:15:00	100.59 100.87	101.13 101.35	100.84 101.09	100.76 101.03	95.375 94.662	95.251 94.53	93.721 97.496	95.14 98.624	93.64 98.065	95.036 99.578	1.1597 1.1639	134.83 130.93	1727.8 1737.4	74.464 74.512	155.33 155.69	11.08 9.7137	48.794 48.794	0.008 0.008	-0.056 -0.056		0 2.114 0 2.114			102.21 102.21			
20 JUN 01 10:30:00	101.16	101.58	101.35	101.3	95.732	95.551	99.495	100.24	99.811	101.2	1.1681	127.03	1737.5	74.559	156.05	9.8022	48.793	800.0	-0.056		0 2.114	4 8.6604	180.68	102.22	2 101.99	9 95.557	158.86 645.3
20 JUN 01 10:45:00 20 JUN 01 11:00:00	101.44 101.73	101.8 102.03	101.61 101.86	101.57 101.84	96.713 97.694	96.485 97.42	100.54 101.01	101.2 101.81	100.91 101.3	102.22 102.78	1.1744 1.1811	123.13 119.23	1722.2 1730.2	74.606 74.654	156.41 156.77	11.495 11.407	48.793 48.792	0.008 0.008	-0.056 -0.056		0 2.114 0 2.11			102.23 102.24			158.87 636.56
20 JUN 01 11:15:00	102.01	102.25	102.12	102.11	98.675	98.355	101.27	101.38	100.88	102.57	1.1879	115.33	1730.3	74.701	157.13	11.319	48.791	0.008	-0.056		0 2.113	9 8.6577	180.68	102.25	102.0	2 95.562	158.87 674.96
<del></del>	EVAPORAT	OR PARAME	TERS, 0031	END-1																							
Thus	T 450 4	T 450 0	T 450 0	T 450 4	T 450 5	T 450.0	T 450 T	T 450 0	T 450 0	T 450 40	D-150-1 evaporator	L-150-1 evaporator	F350-1 evaporator												T 400 0 4	T 400 0 4	T40044 54004
Time	degrees C	legrees C			degrees C	degrees C		T-150-8 degrees C	T-150-9 degrees C	T-150-10 degrees C			steam flow lb/hour	T-336-1C degrees F	scfm	"wcvac	"wcvac	IN WC	IN WC	PD-130-3-1 IN WC	PD-130-4-1 IN WC	PD-130-1 IN WC	degrees F	T-130-1-1 degrees F	T-130-2-1 degrees F	T-130-3-1 degrees F	T-130-4-1 F-130-1 degrees F scfm
20 JUN 01 14:00:00 20 JUN 01 14:15:00	104.46 104.68	104.73 104.96	104.67 104.84	104.69 104.87	97.59 97.722	97.256 97.524	101.41 101.65	102.21 102.44	101.93 102.16	102.93 103.17	1.232 1.2356	125.57 124.93	1727.6 1731.6	75.222 75.269	161.09 161.45	10.753 10.731	48.784 48.783	0.008 0.008	-0.056 -0.056		0 2.112 0 2.11			102.33 102.34			158.91 656.82 158.91 637.19
20 JUN 01 14:30:00 20 JUN 01 14:45:00	104.9 105.11	105.18 105.41	105.01 105.17	105.04 105.22	97.854 97.985	97.659 97.793	101.89 102.13	102.68 102.91	102.39 102.62	103.41 103.65	1.2391 1.2427	125.03 125.31	1730.3 1725.1	75.316 75.364	161.8 162.16	10.709 10.687	48.783 48.782	0.008 0.008	-0.056 -0.056		0 2.111 0 2.111	8 8.6456	180.69	102.35 102.36			
20 JUN 01 15:00:00	105.27	105.58	105.34	105.39	98.117	97.928	102.37	103.15	102.86	103.89	1.2463	126.04	1729.6	75.411	162.52	10.665	48.781	0.008	-0.056		0 2.111	5 8.6438	180.69	102.37	102.17	7 95.586	158.92 651.96
20 JUN 01 15:15:00 20 JUN 01 15:30:00	105.42 105.57	105.72 105.86	105.51 105.67	105.57 105.74	98.249 98.381	98.063 98.198	102.61 102.84	103.38 103.62	103.09 103.32	104.13 104.36	1.2498 1.2534	127.81 129.52	1734 1725.4	75.458 75.506	162.88 163.24	10.643 10.621	48.781 48.78	0.008 0.008	-0.056 -0.056		0 2.111 0 2.111			102.37 102.38			158.92 677.59
20 JUN 01 15:45:00 20 JUN 01 16:00:00	105.71 105.86	106 106.15	105.84 105.95	105.86 105.93	98.579 99.385	98.333 99.003	103.08 103.32	103.85 104.09	103.55 103.79	104.6 104.84	1.257 1.2605	130.94 132.31	1735.8 1712.9	75.553 75.601	163.6 163.96	10.608 10.611	48.779 48.779	0.008 800.0	-0.056 -0.056		0 2.11 0 2.110		180.69 180.7	102.39 102.4			
20 JUN 01 16:15:00 20 JUN 01 16:30:00	106.01 106.15	106.29 106.43	106 106.05	106 106.06	100.04 100.62	99.628 100.12	103.56 103.8	104.32 104.56	104.02 104.25	105.08 105.32	1.2641 1.2677	133.47 134.43	1729.6	75.648	164.32 164.68	10.614	48.778 48.777	0.008	-0.056		0 2.110	7 8.6392		102.41 102.42	102.22	2 95.594	
20 JUN 01 16:45:00	106.3	106.57	106.1	106.13	101.2	100.61	104.04	104.79	104.48	105.56	1.2712	134.47	1730.1 1721.6	75.695 75.743	165.04	10.617 10.62	48.777	800.0 800.0	-0.056 -0.056		0 2.110 0 2.110	4 8.6373	180.7	102.42	102.24	4 95.598	158.94 646.77
20 JUN 01 17:00:00 20 JUN 01 17:15:00	106.44 106.59	106.71 106.85	106.16 106.21	106.19 106.26	101.77 102.11	101.11 101.6	104.28 104.37	105.03 105.12	104.72 104.83	105.76 105.82	1.2748 1.2784	134.1 133.08	1738.5 1734.4	75.79 75.837	165.4 165.76	10.624 10.627	48.776 48.775	0.008 0.008	-0.056 -0.056		0 2.110 0 2.110			102.43 102.44			158.94 645.82 158.95 644.32
20 JUN 01 17:30:00 20 JUN 01 17:45:00	106.74 106.88	106.99 107.13	106.26 106.31	106.33 106.39	101.93 101.74	102.07 101.6	104.42 104.47	105.17	104.83 104.83	105.88 105.94	1.2819 1.2855	131.77 130.49	1732.4 1715.7	75.885 75.932	166.12 166.48	10.63 10.633	48.775 48.774	0.008 0.008	-0.056 -0.056		0 2.109			102.45 102.46			
										,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	112000	100.10	17 10	70.002	100.10	10.000	10.11	0.000	0.000		2.100	0.0000	100	.020			
	EVAPORATO	OR PARAME	TERS, 0031	STRT-2					1-1-1-1-1		D-150-1	L-150-1	F350-1														
Time	T-150-1	T-150-2	T-150-3	T-150-4				T-150-8	T-150-9	T-150-10	evaporator density	evaporator level	evaporator steam flow							PD-130-3-1	PD-130-4-1	PD-130-1		T-130-1-1	T-130-2-1	T-130-3-1	T-130-4-1 F-130-1
Time 21 JUN 01 08:15:00	T-150-1 degrees C 6 61.744	T-150-2 legrees C 62.986	T-150-3 legrees C 68.576	T-150-4 degrees C 69.401	degrees C 43.992	degrees C 43.095	degrees C 37.156	degrees C 37.268	degrees C 37.428	degrees C 38.441	evaporator density Grams/ml 1.1823	evaporator level Inches 136.66	evaporator steam flow lb/hour 1.6159	degrees F 75.869	scfm 166.25	"wcvac 11.564	"wcvac 48.736	IN WC 0.008	IN WC -0.056	IN WC	IN WC 0 2.100	IN WC 6 8.58	degrees F 180.75	degrees F 102.92	degrees F 102.84	degrees F 4 95.698	degrees F scfm 159.14 647.72
Time	T-150-1 degrees C	T-150-2 legrees C	T-150-3 legrees C	<b>T-150-4</b> degrees C	degrees C	degrees C	degrees C	degrees C	degrees C	degrees C	evaporator density Grams/ml	evaporator level Inches	evaporator steam flow lb/hour	degrees F	scfm	"wcvac	"wcvac	IN WC	IN WC	IN WC	IN WC	IN WC 6 8.58 4 8.5791	degrees F 180.75 180.75	degrees F	degrees F 102.84 102.85	degrees F 4 95.698 5 95.699	degrees F scfm 159.14 647.72
Time 21 JUN 01 08:15:00 21 JUN 01 08:30:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00	T-150-1 degrees C 61.744 97.597 99.175 99.923	T-150-2 degrees C 62.986 98.4 100.02 100.38	T-150-3 legrees C 68.576 97.627 98.757 99.826	T-150-4 degrees C 69.401 97.503 99.515 100.27	43.992 48.193 76.222 93.493	degrees C 43.095 48.396 76.637 93.117	degrees C 37.156 37.211 38.235 42.507	degrees C 37.268 37.359 38.954 43,269	degrees C 37.428 37.49 37.552 38.604	degrees C 38.441 38.814 39.187 40.175	evaporator density Grams/ml 1.1823 1.1819 1.1814 1.1809	evaporator level Inches 136.66 136.59 136.52 136.45	evaporator steam flow lb/hour 1.6159 9.6779 701.28 1132.2	degrees F 75.869 75.89 75.91 75.93	scfm 166.25 166.16 166.06 165.97	"wcvac 11.564 11.56 11.555 11.551	"wcvac 48.736 48.735 48.734 48.734	IN WC 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056	IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100	IN WC 6 8.58 4 8.5791 3 8.5782 1 8.5773	degrees F 180.75 180.75 180.75 180.76	degrees F 102.92 102.93 102.94 102.95	degrees F 2 102.84 3 102.85 4 102.86 5 102.87	degrees F 4 95.698 5 95.699 6 95.701 7 95.702	degrees F scfm 159.14 647.72 159.14 654.22 159.15 665.13 159.15 652.82
Time 21 JUN 01 08:15:00 21 JUN 01 08:30:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00 21 JUN 01 09:15:00 21 JUN 01 09:30:00	T-150-1 degrees C 61.744 97.597 99.175 99.923 100.15 100.38	T-150-2 degrees C 62.986 98.4 100.02 100.38 100.74 101.01	T-150-3 legrees C 68.576 97.627 98.757 99.826 100.31 100.52	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69	degrees C 43.992 48.193 76.222 93.493 96.585 97.776	degrees C 43.095 48.396 76.637 93.117 96.137 97.779	degrees C 37.156 37.211 38.235 42.507 50.239 63.102	degrees C 37.268 37.359 38.954 43.269 51.187 63.852	degrees C 37.428 37.49 37.552 38.604 44.778 57.098	degrees C 38.441 38.814 39.187 40.175 46.309 58.478	evaporator density Grams/ml 1.1823 1.1819 1.1814 1.1809 1.17795	evaporator level Inches 136.66 136.59 136.52 136.45 136.37 136.57	evaporator steam flow  Ib/hour	degrees F 75.869 75.89 75.91 75.93 75.951 75.971	scfm 166.25 166.16 166.06 165.97 165.88 165.79	"wcvac 11.564 11.56 11.555 11.551 11.547 11.543	"wcvac 48.736 48.735 48.734 48.734 48.733 48.732	IN WC 0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056	IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.100 0 2.099 0 2.099	IN WC 6 8.58 4 8.5791 3 8.5782 1 8.5773 9 8.5763 8 8.5754	degrees F 180.75 180.75 180.75 180.76 180.76 180.76	degrees F 102.92 102.93 102.94 102.95 102.96	degrees F 102.84 102.85 102.85 102.87 102.87 102.87	degrees F 4 95.698 5 95.699 6 95.701 7 95.702 7 95.704 8 95.706	degrees F scfm 159.14 647.7, 159.14 654.2, 159.15 665.1, 159.15 652.8, 159.15 647.9, 159.16 660.8,
Time  21 JUN 01 08:15:00 21 JUN 01 08:30:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00 21 JUN 01 09:00:00 21 JUN 01 09:30:00 21 JUN 01 09:45:00 21 JUN 01 10:00:00	T-150-1 degrees C 61.744 97.597 99.175 99.923 100.15 100.38 100.6 100.82	T-150-2 legrees C 62.986 98.4 100.02 100.38 100.74 101.01 101.24 101.47	T-150-3 degrees C 68.576 97.627 98.757 99.826 100.31 100.52 100.73 100.93	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 100.9	degrees C 43.992 48.193 76.222 93.493 96.585 97.776 97.595 96.2	43.095 48.396 76.637 93.117 96.137 97.779 97.469 96.063	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278 92.289	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349 93.465	degrees C 37.428 37.49 37.552 38.604 44.778 57.098 75.452 91.434	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601 91.963	evaporator density  Grams/ml 1.1823 1.1819 1.1814 1.1809 1.1795 1.1711 1.1664	evaporator level Inches 136.66 136.59 136.52 136.45 136.37 136.57 136.84 135.9	evaporator steam flow lb/hour 1.6159 9.6779 701.28 1132.2 1406.8 1734.8 1730.4 1727.8	degrees F 75.869 75.89 75.91 75.93 75.951 75.971 75.992 76.011	scfm 166.25 166.16 166.06 165.97 165.88 165.79 165.7	"wevac 11.564 11.56 11.555 11.551 11.547 11.543 11.538 11.534	*wcvac 48.736 48.735 48.734 48.734 48.733 48.732 48.732 48.731	0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056	IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.100 0 2.099 0 2.099 0 2.099 0 2.099 0 2.099	IN WC 6 8.58 4 8.5791 3 8.5782 1 8.5773 9 8.5763 8 8.5754 6 8.5745 5 8.5736	degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76	degrees F 102.92 102.93 102.94 102.95 102.96 102.97 102.98	degrees F 102.84 102.85 102.86 102.87 102.87 102.88 102.88	degrees F 4 95.698 5 95.699 6 95.701 7 95.702 7 95.704 8 95.706 9 95.707 9 95.709	degrees F scfm 159.14 647.7: 159.14 654.2: 159.15 665.1: 159.15 62.8: 159.16 660.8: 159.16 644.4: 159.16 644.3:
Time 21 JUN 01 08:15:00 21 JUN 01 08:30:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00 21 JUN 01 09:15:00 21 JUN 01 09:30:00 21 JUN 01 09:45:00	T-150-1 degrees C 6 61.744 97.597 99.175 99.923 100.15 100.38 100.6	T-150-2 degrees C 62.986 98.4 100.02 100.38 100.74 101.01 101.24	T-150-3 legrees C 68.576 97.627 98.757 99.826 100.31 100.52 100.73	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 100.9	degrees C 43.992 48.193 76.222 93.493 96.585 97.776 97.595	degrees C 43.095 48.396 76.637 93.117 96.137 97.779 97.469	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349	degrees C 37.428 37.49 37.552 38.604 44.778 57.098 75.452	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601	evaporator density Grams/mi 1.1823 1.1819 1.1814 1.1809 1.1795 1.1711 1.1664	evaporator level Inches 136.66 136.59 136.52 136.45 136.37 136.57 136.84	evaporator steam flow lb/hour 1.6159 9.6779 701.28 1132.2 1406.8 1734.8 1730.4	degrees F 75.869 75.89 75.91 75.93 75.951 75.971 75.992	scfm 166.25 166.16 166.06 165.97 165.88 165.79	"wcvac 11.564 11.56 11.555 11.551 11.547 11.543 11.538	"wcvac 48.736 48.735 48.734 48.734 48.733 48.732 48.732	IN WC 0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056 -0.056	IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.100 0 2.100 0 2.099 0 2.099 0 2.099	IN WC 6 8.58 4 8.5791 3 8.5782 1 8.5773 8 8.5763 8 8.5745 5 8.5736 3 8.5727	degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76 180.76	degrees F 102.92 102.93 102.94 102.95 102.96 102.96	degrees F 102.84 102.85 102.86 102.87 102.87 102.88 102.88 102.9	degrees F 4 95.698 5 95.699 6 95.701 7 95.702 7 95.704 8 95.706 9 95.707 9 95.707	degrees F scfm 159.14 647.7; 159.14 654.2; 159.15 665.1; 159.15 652.8; 159.16 647.9; 159.16 644.4; 159.16 642.3; 159.17 662.7;
Time  21 JUN 01 08:15:00 21 JUN 01 08:35:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00 21 JUN 01 09:30:00 21 JUN 01 09:30:00 21 JUN 01 09:30:00 21 JUN 01 10:00:00 21 JUN 01 10:15:00 21 JUN 01 10:30:00 21 JUN 01 10:30:00 21 JUN 01 10:45:00	T-150-1 degrees C 61.744 97.597 99.175 99.923 100.15 100.38 100.6 100.82 101.02 101.21	T-150-2 degrees C 62.986 98.4 100.02 100.38 100.74 101.01 101.24 101.47 101.71 101.95 102.19	T-150-3 legrees C 68.576 97.627 98.757 99.826 100.31 100.52 100.73 100.93 101.22 101.71 102.19	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 101.11 101.51	degrees C 43.992 48.193 76.222 93.493 96.585 97.776 97.595 96.2 94.211	degrees C 43.095 48.396 76.637 93.117 96.137 97.779 97.469 96.063 94.088 95.2 96.137	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278 92.289 96.806 98.98 100.07	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349 93.465 97.873 99.864 101.01	degrees C 37.428 37.49 37.552 38.604 44.778 57.098 75.452 91.434 97.301 99.488 100.62	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601 91.963 98.899 100.89	evaporator density  Grams/ml  1.1823 1.1819 1.1814 1.1809 1.1795 1.1711 1.1664 1.1624 1.1648 1.1695 1.1739	evaporator level     136.66   136.59   136.52   136.45   136.37   136.57   136.84   135.9   132.32   128.36   124.34	evaporator steam flow Ib/hour 1.6159 9.6779 701.28 1132.2 1406.8 1734.8 1730.4 1727.8 1729.9 1729.8	degrees F 75.869 75.89 75.91 75.93 75.951 75.971 75.992 76.011 76.032 76.052 76.072	scfm 166.25 166.16 166.06 165.97 165.88 165.79 165.7 165.61 165.52 165.43	"wevac 11.564 11.565 11.555 11.551 11.547 11.543 11.538 11.534 11.53 11.526 11.521	**Wevac 48.736 48.735 48.734 48.733 48.732 48.731 48.73 48.73 48.73 48.73 48.73	0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056 -0.056	IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.190 0 2.099 0 2.099 0 2.099 0 2.099 0 2.099 0 2.099 0 2.099	IN WC 8.58 8.5791 3 8.5782 1 8.5773 9 8.5754 6 8.5745 3 8.5727 2 8.5717 9 8.5708	degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76 180.76 180.76	degrees F 102.92 102.93 102.94 102.95 102.96 102.97 102.98 103.99 103.90	degrees F 102.88 102.86 102.87 102.87 102.87 102.88 102.89 102.99 102.99	degrees F 4 95.698 5 95.699 6 95.701 7 95.702 8 95.706 9 95.707 9 95.709 1 95.711 2 95.712 3 95.714	degrees F scfm 159.14 647.7; 159.14 654.2; 159.15 665.1; 159.15 647.9; 159.16 660.8; 159.16 644.4; 159.16 642.3; 159.17 662.7; 159.17 641.5;
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Time  21 JUN 01 08:15:00 21 JUN 01 08:30:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00 21 JUN 01 09:30:00 21 JUN 01 09:45:00 21 JUN 01 09:45:00 21 JUN 01 10:30:00 21 JUN 01 10:15:00 21 JUN 01 10:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:30:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00	T-150-1  degrees C (61.744 97.597 99.175 99.923 100.15 100.38 100.6 100.82 101.02 101.21 101.4 101.6 101.79 101.98 102.37  EVAPORATO  T-150-1  degrees C C 103.53 103.72	T-150-2 legrees C 62.986 98.4 100.02 100.38 100.74 101.01 101.24 101.47 101.71 101.95 102.19 102.42 102.66 102.79 102.92 103.05	T-150-3 legrees C 68.576 97.627 99.757 99.826 100.31 100.52 100.73 101.22 101.71 102.19 102.68 102.84 102.94 103.04 103.14  TERS, 0031	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 100.9 101.11 101.51 101.97 102.44 102.9 103.18 103.26  END-2  T-150-4 degrees C 103.79 103.79	degrees C 43,992 48,193 76,222 93,493 96,585 97,776 97,595 96,22 94,211 95,327 96,277 98,162 98,923 95,33	43.095 48.396 76.637 93.117 96.137 97.779 97.469 96.063 94.088 95.2 96.137 97.075 98.013 98.691 95.229	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278 92.289 96.806 98.98 100.02 100.82 99.07 77.764 98.296	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349 93.465 97.873 99.864 101.01 101.64 101.59 98.815 98.835 99.176	degrees C 37.428 37.49 37.552 38.604 44.778 57.098 75.452 91.434 97.301 99.488 100.62 101.25 100.85 99.27 98.483 98.954	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601 91.963 98.899 100.89 102.04 102.56 102.89 99.51 99.779 100.05	evaporator density  Grams/mi  1.1823 1.1819 1.1814 1.1809 1.1795 1.1711 1.1664 1.1624 1.1624 1.1685 1.1739 1.1781 1.1824 1.1867 1.1909 1.1952  D-150-1 evaporator density  Grams/mi 1.2209 1.2251	evaporator level Inches 136.66 136.59 136.52 136.45 136.37 136.57 136.84 135.99 132.32 128.36 124.34 120.3 116.27 121.79 124.49 126.51  L-150-1 evaporator level nches 135.91 133.45	evaporator steam flow Ib/hour 1.6159 9.6779 701.28 1132.2 1406.8 1730.4 1727.8 1729.9 1729.9 1729.9 1724.1 1746.7 1715.2	degrees F 75.869 75.89 75.91 75.93 75.951 75.971 75.992 76.011 76.032 76.052 76.072 76.093 76.113 76.133 76.154 76.174	scfm 166.25 166.16 166.06 165.97 165.79 165.7 165.61 165.52 165.43 165.25 165.48 165.26 164.97 164.88	"wcvac 11.564 11.555 11.555 11.551 11.543 11.538 11.526 11.527 11.543 11.537 11.526 11.527 11.513 11.509 11.504 11.5	**Wcvac 48.736 48.735 48.731 48.732 48.732 48.731 48.73 48.73 48.729 48.729 48.729 48.726 ***Page 48.726 ***Page 48.727 48.727 48.726 ***Page 48.727 48.727 48.726 ***Page 48.727 48.727 48.726 ***Page 48.727 48.727 48.726 ***Page 48.727 48.727 48.727 48.726 ***Page 48.727 48.727 48.727 48.726 ***Page 48.727 48.727 48.726 ***Page 48.727 4	IN WC  0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008 0.008	IN WC -0.056	IN WC  PD-130-3-1 IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.100 0 2.099 0 2.099 0 2.099 0 2.099 0 2.099 0 2.098 0 2.098 0 2.098 0 2.098 0 1.098	IN WC 8.58 8.5791 8.57793 8.5763 8.5754 6.8.5736 3.8.5722 8.5717 9.8.5708 8.5669 7.8.5662 PD-130-1 IN WC 3.5662	degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.77	degrees F 102.92 102.93 102.94 102.95 102.96 102.97 103.93 103.01 103.02 103.03 103.04	degrees F 102.84 102.85 102.86 102.86 102.86 102.86 102.86 102.86 102.86 102.86 102.86 102.86 102.96	degrees F 4 95.698 5 95.699 6 95.701 7 95.702 8 95.706 9 95.707 9 95.709 1 95.711 2 95.712 3 95.714 4 95.715 5 95.717 6 95.719 7 95.722 7 95.722	degrees F         scfm           159.14         647.77           159.14         654.22           159.15         655.28           159.15         652.83           159.15         647.93           159.16         660.81           159.16         644.41           159.17         642.33           159.17         641.5           159.18         662.74           159.18         664.00           159.19         644.02           159.19         644.02           159.19         647.82    T-130-4-1  degrees F  scfm  559.21  640.72  159.21  640.72  640.72  159.21  640.72
Time  21 JUN 01 08:15:00 21 JUN 01 08:30:00 21 JUN 01 08:45:00 21 JUN 01 09:00:00 21 JUN 01 09:45:00 21 JUN 01 09:45:00 21 JUN 01 09:45:00 21 JUN 01 10:15:00 21 JUN 01 10:15:00 21 JUN 01 10:30:00 21 JUN 01 11:00:00 21 JUN 01 11:30:00 21 JUN 01 11:15:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00 21 JUN 01 11:45:00	T-150-1  degrees C c 61.744 97.597 99.175 99.923 100.6 100.82 101.02 101.02 101.04 101.6 101.79 101.98 102.18 102.37  EVAPORATO  T-150-1  degrees C c 103.53 103.72 103.91 104.11	T-150-2 legrees C	T-150-3 legrees C 68.576 97.627 98.757 99.826 100.31 100.52 100.73 100.93 101.22 101.71 102.19 102.68 102.84 103.04 103.14  TERS, 0031I  T-150-3 legrees C 103.9 104.16 104.43 104.7	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 100.9 101.11 101.51 101.97 102.94 103 103.09 103.18 103.26  END-2  T-150-4 degrees C 103.79 103.96 104.24 104.52	degrees C 43,992 48,193 76,222 93,493 96,595 97,776 97,595 96,272 97,217 98,162 98,228 95,13 95,32 7-150-5 degrees C 97,637 98,124 98,612 98,612 98,98	degrees C 43.095 48.396 76.637 93.117 97.779 97.469 96.063 94.088 95.2 96.137 97.075 98.013 98.691 95.048 95.229	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278 92.289 96.806 98.98 100.07 100.82 100.82 99.67 97.764 98.296	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349 93.465 97.873 99.864 101.01 101.64 101.59 98.815 99.176	7.150-9 degrees C 37.428 37.49 37.552 38.604 44.778 57.098 75.452 91.434 97.301 99.488 100.62 101.25 100.85 99.27 98.483 98.954	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601 91.963 98.899 100.89 102.56 102.59 100.05  T-150-10 degrees C 102.58 102.94 103.55	evaporator density  Grams/ml  1.1823 1.1819 1.1814 1.1809 1.1795 1.1711 1.1664 1.1624 1.1648 1.1695 1.1739 1.1781 1.1827 1.1909 1.1952  D-150-1 evaporator density  Grams/ml 1.2209 1.2251 1.2294 1.2337	evaporator   level	evaporator   steam flow    b/hour   1.6159   9.6779     701.28   1132.2     1406.8   1734.8     1730.4   1727.8     1729.9   1729.9     1724.1     1746.7     1715.2     F350-1     evaporator   steam flow    b/hour   1728.3     1733.5     1730.5     1727.2	degrees F 75.869 75.89 75.91 75.93 75.951 75.971 75.992 76.011 76.032 76.052 76.072 76.093 76.113 76.133 76.154 76.174	scfm 166.25 166.16 166.06 165.97 165.79 165.79 165.52 165.43 165.34 165.25 165.15 165.06 164.97 164.88	"wcvac 11.564 11.555 11.555 11.551 11.543 11.538 11.534 11.53 11.526 11.521 11.517 11.513 11.504 11.5 11.504 11.5 11.504 11.5 11.5 11.5 11.5 11.5 11.5 11.5 11.	**Wcvac 48.736 48.736 48.735 48.731 48.732 48.731 48.73 48.729 48.729 48.727 48.727 48.726 ***Provided Research 130-2 ***Provided	IN WC  0.008	IN WC -0.056	IN WC  PD-130-3-1 IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.099 0 2.099 0 2.099 0 2.099 0 2.099 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098	IN WC 8.58 8.5791 8.5792 8.5763 8.5754 6.5736 8.5736 8.5736 8.5736 8.5736 8.5736 8.5736 8.5736 8.5736 8.5736 8.55699 7.8.5662 PD-130-1 IN WC 8.5607 1.8.5558 8.5559 8.5558 8.5559 8.5558 8.5558 8.5559 8.5558 8.5559 8.5558 8.5559 8.5558 8.5559 8.5558 8.5558 8.5559	degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.77 180.77	degrees F 102.92 102.93 102.94 102.95 102.96 102.97 103.93 103.01 103.02 103.04 103.04	degrees F	degrees F 4 95.698 5 95.698 6 95.701 7 95.702 8 95.706 9 95.707 9 95.709 1 95.711 2 95.712 2 95.713 3 95.714 4 95.715 5 95.717 6 95.722  T-130-3-1 degrees F 4 95.731 5 95.733 5 95.733 7 95.736	degrees F scfm 159.14 647.7; 159.15 652.8; 159.15 652.8; 159.16 660.8; 159.16 644.4; 159.17 641.5; 159.17 632.2; 159.18 662.7; 159.17 633.2; 159.18 664.0; 159.18 642.9; 159.19 647.8;  T-130-4-1 degrees F scfm 159.21 649.2; 159.21 649.2; 159.21 649.2; 159.21 649.2; 159.21 649.2;
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159.15 652.8; 159.15 652.8; 159.15 647.9; 159.16 644.9; 159.17 642.3; 159.17 642.3; 159.17 641.1; 159.17 633.2; 159.18 662.7; 159.18 664.0; 159.18 642.9; 159.19 647.8;  T-130-4-1 F-130-1 degrees F scfm 159.21 649.2; 159.21 649.2; 159.22 637.7; 159.22 637.7; 159.23 659.5; 159.23 659.5; 159.23 665.6; 159.23 674.3; 159.23 674.3; 159.23 674.3; 159.23 674.3; 159.23 674.3; 159.23 674.3; 159.24 670.66
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101.71 101.95 102.19 102.42 102.66 102.79 102.92 103.05  PARAME  T-150-2 legrees C c 103.88 104.12 104.35 104.35 104.55 104.59 104.83 105.07 105.54 105.78 105.94 106.08 106.22 106.39	T-150-3 legrees C 68.576 97.627 99.826 100.31 100.52 100.73 100.93 101.22 101.71 102.19 102.88 102.84 103.04 103.14  T-150-3 legrees C 103.9 104.16 104.43 104.97 105.52 105.68 105.84 106.15 106.15 106.15 106.17	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 100.9 101.11 101.51 101.97 102.44 102.9 103.18 103.26  END-2  T-150-4 degrees C 103.79 104.84 105.51 105.66 105.84 106.01 106.18 106.36 106.53	degrees C 43,992 48,193 76,222 93,493 96,585 97,776 97,595 96,22 94,211 95,327 96,272 97,217 98,162 98,928 95,13 95,32   T-150-5 degrees C 69,693 99,49 99,319 99,49 99,149 99,319 99,49 100,36 100,63	degrees C 43.095 48.396 76.637 93.117 96.137 97.779 97.469 96.063 94.088 95.22 96.137 97.075 98.013 98.691 95.229 97.437 97.97.99 98.381 98.784 98.942 99.099 99.256 99.414 99.571 99.692 99.756 99.82 99.884 99.949	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278 92.289 96.806 98.98 100.07 7100.82 190.82 99.06 7100.82 99.06 7100.82 100.82 99.06 100.07 97.764 98.296 100.07 97.764 100.07 97.764 100.07 97.764 100.07 97.764 100.07 97.764 98.296 100.07 97.764 98.296 100.07 97.764 98.296 100.07 97.764 98.296 100.07 97.764 98.296 100.07 97.764 98.296 100.07 97.764 100.07	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349 93.465 97.873 99.864 101.01 101.64 101.59 98.815 99.176  T-150-8 degrees C 102.03 102.46 102.65 102.84 103.03 103.22 103.38 103.55 103.62 103.74 103.86 103.98 104.1 104.22	7.150-9 degrees C 37.428 37.439 37.552 38.604 44.778 57.098 75.452 91.434 97.301 99.488 100.62 101.25 100.85 99.27 98.483 98.954  7.150-9 degrees C 101.27 101.66 101.94 102.69 102.69 102.85 103.01 103.17 103.33 103.48 103.84 103.82	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601 91.963 98.899 100.89 102.56 102.89 99.51 99.779 100.05  T-150-10 degrees C 102.58 102.94 103.25 103.25 103.55 104.45 104.63 104.79 104.95 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.26 105.41	evaporator   density	evaporator   level     Inches   136.66   136.59   136.52   136.45   136.37   136.57   136.84   135.9   132.32   128.36   124.34   120.3   116.27   121.79   124.49   126.51     L-150-1   evaporator   level   133.45   132.47   132.23   131.99   131.62   131.11   130.6   129.82   128.93   128.78   129.03   129.28   131.12	evaporator steam flow   Ib/hour	degrees F 75.869 75.89 75.991 75.993 75.951 75.971 75.992 76.011 76.032 76.052 76.072 76.093 76.113 76.133 76.154 76.174   T-336-1C degrees F 76.296 76.316 76.357 76.387 76.387 76.387 76.499 76.499 76.499 76.542 76.54	scfm 166.25 166.16 166.06 165.97 165.79 165.77 165.61 165.52 165.43 165.25 165.15 165.06 164.97 164.88	"wcvac 11.564 11.555 11.555 11.551 11.543 11.533 11.526 11.527 11.543 11.539 11.574 11.573 11.504 11.51 11.504 11.51 11.504 11.51 11.504 11.504 11.51 11.504 11.51 11.504 11.51 11.504 11.475 1	**Wevac 48.736 48.736 48.735 48.731 48.732 48.732 48.731 48.73 48.73 48.729 48.729 48.727 48.726 ***  **I30-2 PC ***  **Wevac 48.722 48.721 48.721 48.721 48.721 48.717 48.717 48.717 48.715 48.715 48.715 48.715 48.715 48.713	IN WC  0.008	IN WC -0.056	PD-130-3-1 IN WC	PD-130-4-1 IN WC  PD-130-4-1 IN WC  2.099  PD-130-4-1 IN WC  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.099  2.098  2.096  2.096  2.096  2.096  2.096  2.096  2.096  2.096  2.096  2.096  2.096  2.096  2.096	IN WC 8.58 8.5791 8.5792 8.5763 8.5754 8.5662 PD-130-1 IN WC 3.5597 8.5662 PD-130-1 8.5597 8.5598 8.5593 8.5551 8.5552 8.5551 8.5542 8.55542 8	T-335-2  degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.77 180.78	degrees F 102.92 102.93 102.94 102.95 102.96 102.97 102.98 103.01 103.01 103.02 103.03 103.04 103.04  T-130-1-1 degrees F 103.09 103.11 103.12 103.13 103.14 103.15 103.16 103.17 103.18 103.19 103.19	degrees F 102.94 103.04 103.05 103.05 103.01 103.15 103.16	degrees F 4 95.698 5 95.698 6 95.701 7 95.702 8 95.706 9 95.707 9 95.707 9 95.712 2 95.712 3 95.714 4 95.715 5 95.717 6 95.722  T-130-3-1 degrees F 95.731 95.733 95.734 95.735 7 95.736 8 95.737 95.736 95.737 95.736 95.737 95.738 95.744 95.743 95.745 95.746 95.748 95.748 95.748 95.748	degrees F scfm 159.14 647.7; 159.15 652.8; 159.15 665.1; 159.15 652.8; 159.16 660.8; 159.16 644.2; 159.17 641.4; 159.17 632.2; 159.18 662.7; 159.18 664.0; 159.18 664.0; 159.18 644.9; 159.19 647.8;  T-130-4-1 fedgrees F scfm 159.21 649.2; 159.22 637.7; 159.22 637.7; 159.23 659.5; 159.23 659.5; 159.23 665.68; 159.23 674.3; 159.24 670.6; 159.24 638.04; 159.24 661.15; 159.24 661.15; 159.25 673.6; 159.25 673.6; 159.26 673.6; 159.27 673.6; 159.28 665.66; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.29 673.6; 159.25 673.6; 159.25 673.6;
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102.66 102.79 102.92 103.05  T-150-2 legrees C (30.88 104.12 104.35 104.59 104.83 105.07 105.31 105.54 105.78 105.94 106.08 106.22 106.35	T-150-3 legrees C 68.576 97.627 98.757 99.826 100.31 100.52 100.73 100.93 101.22 101.71 102.19 102.68 102.94 103.04 103.14  TERS, 00311  T-150-3 legrees C 103.9 104.13 104.7 104.97 105.21 105.37 105.52 105.68 106.15 106.31	T-150-4 degrees C 69.401 97.503 99.515 100.27 100.48 100.69 100.9 101.11 101.51 101.97 102.94 103.18 103.26  END-2  T-150-4 degrees C 103.79 103.96 104.24 104.52 104.8 105.07 105.31 105.49 105.64 106.18 106.01 106.18 106.03	degrees C 43,992 48,193 76,222 93,493 96,595 96,272 94,211 95,327 96,272 97,217 98,162 98,928 95,13 95,32 77,637 98,124 98,612 98,98 99,149 99,319 99,488 99,637 99,827 100,09 100,36 100,63 100,09	degrees C 43.095 48.396 76.637 93.117 97.779 97.469 96.063 94.088 95.2 96.137 97.075 98.013 98.691 95.048 95.229   T-150-6 degrees C 697.437 97.909 98.381 98.784 98.942 99.099 99.256 99.414 99.571 99.692 99.756 99.884	degrees C 37.156 37.211 38.235 42.507 50.239 63.102 81.278 92.289 96.806 98.98 100.07 100.82 100.82 99.067 97.764 98.296   T-150-7 degrees C 101.1 101.71 101.99 102.28 102.46 102.61 102.67 102.79 102.86 102.98	degrees C 37.268 37.359 38.954 43.269 51.187 63.852 82.349 93.465 97.873 99.864 101.01 101.64 101.59 98.815 99.176 T-150-8 degrees C 102.03 102.46 102.65 102.84 103.03 103.03 103.52 103.38 103.52 103.74 103.98 103.98	7-150-9 degrees C 101.27 102.49 102.49 102.69 103.8 10	degrees C 38.441 38.814 39.187 40.175 46.309 58.478 76.601 91.963 98.899 100.89 102.04 102.56 102.89 99.51 99.779 100.05  T-150-10 degrees C 102.58 102.94 103.25 103.85 104.15 104.46 104.63 104.79 104.95 105.41	evaporator density  Grams/ml  1.1823 1.1819 1.1814 1.1809 1.1795 1.1711 1.1664 1.1624 1.1648 1.1695 1.1739 1.1781 1.1827 1.1909 1.1952  D-150-1 evaporator density  Grams/ml 1.2209 1.2251 1.2337 1.238 1.2422 1.2465 1.2508 1.2553 1.2593 1.2636 1.2679 1.2721	evaporator   level	evaporator steam flow  bb/hour	75.869 75.89 75.991 75.991 75.993 75.951 75.971 75.992 76.011 76.032 76.052 76.072 76.093 76.113 76.133 76.154 76.174   T-336-1C degrees F 76.296 76.316 76.337 76.357 76.377 76.398 76.418 76.438 76.459 76.499 76.592 76.554	scfm 166.25 166.16 166.06 165.97 165.79 165.79 165.52 165.43 165.34 165.25 165.15 165.06 164.97 164.88 F 164.33 164.24 164.15 164.06 163.97 163.88 163.78 163.69 163.61 163.42 163.33 163.24	"wcvac 11.564 11.555 11.555 11.551 11.551 11.543 11.538 11.534 11.53 11.526 11.527 11.517 11.513 11.504 11.51 11.504 11.51 11.504 11.475 11.476 11.462 11.488 11.454 11.445 11.445 11.445 11.445 11.445 11.445 11.445 11.445 11.445 11.442 11.424 11.424 11.424 11.424	**Wcvac 48.736 48.736 48.735 48.731 48.732 48.732 48.731 48.73 48.729 48.729 48.727 48.726 ***  **I30-2** **Wcvac 48.727 48.721 48.721 48.721 48.721 48.721 48.721 48.721 48.719 48.719 48.719 48.719 48.717 48.716 48.715 48.715 48.715 48.715	IN WC  0.008	IN WC -0.056	PD-130-3-1 IN WC	IN WC 0 2.100 0 2.100 0 2.100 0 2.100 0 2.100 0 2.099 0 2.099 0 2.099 0 2.099 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.098 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.096 0 2.095 0 2.095	IN WC 8.58 4 8.5791 3 8.5782 1 8.5773 9 8.568 8 8.5746 6 8.5746 6 8.5746 6 8.5746 6 8.5746 6 8.5746 6 8.5746 7 8.569 8 8.6699 7 8.569 8 8.6602  PD-130-1 IN WC 3 8.5602  PD-130-1 IN WC 3 8.5602  PD-130-1 IN WC 3 8.56062  SSE 8.5662  PD-130-1 IN WC 3 8.56062  RSE 8.5662  PD-130-1 IN WC 3 8.56062  RSE 8.5662  RSE 8.5662  RSE 8.5663 RSE 8.5563 RSE 8.5663	T-335-2 degrees F 180.75 180.75 180.75 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.76 180.77 180.77 180.77 180.77 180.77 180.77 180.78 180.78 180.78 180.78 180.78 180.78 180.78 180.78 180.78 180.78 180.78	degrees F 102.92 102.93 102.94 102.95 102.96 102.97 103.93 103.01 103.02 103.03 103.04 103.04 103.04 103.03 103.01 103.03 103.01 103.03 103.11 103.12 103.12 103.13 103.14 103.15 103.16	degrees F 102.96 102.87 102.87 102.87 102.87 102.87 102.87 102.97	degrees F 4 95.698 5 95.698 6 95.701 7 95.702 8 95.706 9 95.707 9 95.709 1 95.711 2 95.712 3 95.714 4 95.715 5 95.717 6 95.722  T-130-3-1 degrees F 4 95.731 95.733 6 95.735 7 95.738 9 95.744 1 95.743 1 95.744 1 95.744 1 95.745 2 95.744 3 95.746 4 95.748 5 95.746 6 95.752 7 95.754	degrees F scfm 159.14 647.77 159.14 654.21 159.15 652.81 159.15 647.93 159.16 660.81 159.16 642.33 159.17 642.33 159.17 642.33 159.17 641.59.18 662.74 159.18 664.00 159.18 664.00 159.18 642.33 159.19 647.82 159.19 647.82 159.19 647.82 159.19 647.82 159.21 640.72 159.22 636.57 159.22 636.57 159.23 659.58 159.23 659.58 159.23 659.58 159.23 674.34 159.24 661.13 159.25 673.85 159.2



# **ENGINEERING DESIGN FILE**

EDF- 2506 Rev. No. 0

# NWCF Fluoride Hot Sump Tank - NCC-119

# Metals, Anions, and Miscellaneous

Analyte	Units	Method Number	Sample L 981203		Sample L 990226		Sample Le 990306		Sample L 990317	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
рН		EPA150.1	0.45		0.35	В	0.34	U	0.34	U
Acidity	N	AC7012								
Aluminum	μg/L	SW6010B			8.29E+06		3.04E+07		2.31E+07	
Antimony	μg/L	SW6010B	4.52E+02	U	4.40E+02	U	2.82E+03	U	2.82E+03	U
Arsenic	μg/L	SW6010B	5.04E+02	U	4.74E+02	U	5.74E+03	UN	5.74E+03	UN
Barium	μg/L	SW6010B	1.11E+03		2.94E+02		1.80E+03	В	1.53E+03	В
Beryllium	μg/L	SW6010B	1.8E+01	В	3.8E+01	В	1.31E+02	В	1.11E+02	В
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	6.59E+04		3.85E+04		1.43E+05		1.06E+05	
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171								
Chromium	μg/L	SW6010B	2.64E+04		3.34E+04		1.07E+05		9.03E+04	
Cobalt	μg/L	SW6010B			2.74E+02	В .	1.25E+03	В	1.64E+03	В
Copper	μg/L	SW6010B			2.75E+03		5.40E+03	Ε	5.80E+03	Е
Fluoride	µg/L	AC7093	5.77E+05	В	8.84E+05	ΒE	2.45E+06	ΒE	2.80E+06	В
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	1.82E+04		1.35E+04		4.95E+04	N	3.83E+04	N
Manganese	μg/L	SW6010B			3.78E+04		1.42E+05		1.13E+05	
Mercury	μg/L	SW7470A	1.19E+05		1.95E+07	N	1.41E+07		1.58E+07	
Nickel	μg/L	SW6010B	1.19E+04		1.15E+04	N	2.73E+04		2.20E+04	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	3.32E+02	U	5.34E+02	U	3.14E+03	U	3.14E+03	U
Silver	µg/L	SW6010B	1.38E+02	В	3.18E+02	В	2.32E+02	U	2.32E+02	U
Sodium	μg/L	SW6010B								
Sulfur	µg/L	SW6010B								
Thallium	μg/L	SW6010B	4.68E+02	U	5.68E+02	U	3.76E+03	U	3.76E+03	U
Uranium	μg/L	AC7920			5.41E+03		1.46E+04		9.68E+03	
Vanadium	μg/L	SW6010B	1.64E+02	U	1.12E+02	В	4.54E+02	U	4.54E+02	U
Zinc	μg/L	SW6010B	2.01E+04		8.36E+03		1.42E+04	Е	1.20E+04	Е
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972								
TIC	μg/L	AC8060			4.66E+04	U	5.82E+04	U	5.82E+04	U
TOC	μg/L	SW9060			1.29E+05	В	3.06E+05	В	3.06E+05	В

#### **ENGINEERING DESIGN FILE**

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# NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Metals, Anions, and Miscellaneous (con't)

Analyte	Units	Method Number	Sample L 990320	_	Sample L 990407	-	Sample L 990411		Sample L 990624	_
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1	0.34	U	0.34	U	0.34	U		
Acidity	N	AC7012							5.8E00	
Aluminum	μg/L	SW6010B	2.16E+07		3.20E+07		2.55E+07		2.05E+07	
Antimony	μg/L	SW6010B	2.82E+03	U	2.82E+03	U	2.82E+03	U	7.60E+02	U
Arsenic	μg/L	SW6010B	9.19E+03	BN	5.74E+03	U	5.74E+03	Ü	2.59E+03	В
Barium	μg/L	SW6010B	1.55E+03	В	1.94E+03	В	2.00E+03	В	7.66E+02	E
Beryllium	μg/L	SW6010B	1.01E+02	В	1.72E+02	В	1.31E+02	В	1.01E+02	
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	1.09E+05		3.70E+04	Ε	2.90E+04	E	1.46E+04	
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171								
Chromium	μg/L	SW6010B	9.73E+04		9.67E+04		9.97E+04		4.51E+04	
Cobalt	μg/L	SW6010B	3.74E+02	U	4.85E+02	В	5.45E+02	В	3.15E+02	В
Copper	μg/L	SW6010B	5.43E+03	Е	6.73E+03	Ε	6.53E+03	E	3.45E+03	
Fluoride	μg/L	AC7093	3.38E+06		1.94E+06		2.10E+06	В	6.95E+05	В
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	3.82E+04	N	1.02E+05	E	9.68E+04	Е	4.55E+04	
Manganese	μg/L	SW6010B	1.07E+05		2.12E+05		2.03E+05		1.02E+05	
Mercury	μg/L	SW7470A	2.19E+07		2.06E+07		2.13E+07		2.97E+06	
Nickel	μg/L	SW6010B	2.25E+04		2.28E+04		2.17E+04		1.40E+04	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	µg/L	SW6010B								
Selenium	μg/L	SW6010B	3.14E+03	U	3.14E+03	U	3.14E+03	U	6.12E+02	U
Silver	μg/L	SW6010B	2.32E+02	U	2.32E+02	U	2.32E+02	U	9.9E+01	В
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	3.76E+03	U	3.76E+03	U	3.76E+03	U	1.11E+03	U
Uranium	μg/L	AC7920	1.02E+04		1.65E+04		1.37E+04		8.49E+03	
Vanadium	μg/L	SW6010B	4.54E+02	U	4.54E+02	U	4.54E+02	U	1.71E+02	В
Zinc	μg/L	SW6010B	1.01E+04	E	1.68E+04	E	1.48E+04	E	9.45E+03	
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972								
TIC	μg/L	AC8060	5.82E+04	U	5.82E+04	U	5.82E+04	U	6.83E+04	U
TOC	μg/L	SW9060	3.91E+05	В	1.19E+05		7.54E+04	В	1.92E+05	В

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# NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Metals, Anions, and Miscellaneous (con't)

Analyte	Units	Method Number	Sample Log # 9909071		Sample Log # 9910191		Sample Log # 9911071		Sample Log # 0001121	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1								
Acidity	N	AC7012	1.6E00		4.3E-01	В	1.4E00		1.1E00	
Aluminum	μg/L	SW6010B	2.42E+06		1.49E+06		1.67E+06		1.53E+06	
Antimony	µg/L	SW6010B	7.56E+02	U	7.56E+02	U	7.56E+02	U	1.58E+02	U
Arsenic	μg/L	SW6010B	8.34E+02	U	2.57E+03	В	1.14E+03	В	5.17E+02	Ų
Barium	μg/L	SW6010B	2.69E+02	ΒE	7.98E+01	ΒE	1.91E+02	ΒE	2.23E+02	
Beryllium	μg/L	SW6010B	1.5E+01	В	< 4.2E00	U	6.3E00	В	9E00	В
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	4.41E+03		1.09E+03		2.95E+03		2.54E+03	
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171								
Chromium	μg/L	SW6010B	6.66E+03		2.96E+03		7.76E+03		6.12E+03	Ε
Cobalt	μg/L	SW6010B	7.98E+01	U	7.98E+01	U	1.16E+02	В	9.8E+01	В
Copper	μg/L	SW6010B	8.67E+02		2.50E+02	В	7.73E+02		9.61E+02	Ε
Fluoride	μg/L	AC7093	6.01E+04	В	2.42E+04	В	7.30E+04	В	6.69E+04	В
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	5.51E+03		2.24E+03	В	4.53E+03		4.10E+03	
Manganese	μg/L	SW6010B	1.78E+04		6.98E+03		1.65E+04		1.38E+04	
Mercury	μg/L	SW7470A	2.51E+03	Ε	9.28E+02	ΒE	3.42E+03		2.83E+03	ВЕ
Nickel	μg/L	SW6010B	5.72E+03		1.98E+03		5.42E+03		4.20E+03	
Nitrate	µg/L	AC7074								
Phosphorus	µg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	6.09E+02	U	6.09E+02	U	6.09E+02	U	8.76E+02	U
Silver	μg/L	SW6010B	4.4E+01	U	4.4E+01	Ų	4.4E+01	U	6.6E+01	В
Sodium	µg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	1.11E+03	U	1.11E+03	U	1.11E+03	U	4.87E+02	U
Uranium	µg/L	AC7920	1.51E+03		6.99E+02		1.03E+03		7.33E+02	
Vanadium	µg/L	SW6010B	7.77E+01	U	7.77E+01	U	7.77E+01	U	3.5E+01	U
Zinc	µg/L	SW6010B	1.85E+03		6.11E+02		1.60E+03		1.49E+03	
Zirconium	µg/L	SW6010B								
UDS	μg/L	AC7972								
TIC	μg/L	AC8060	2.70E+03	UE	3.45E+03	U	3.45E+03	U	2.70E+04	U
тос	µg/L	SW9060	4.81E+04		1.32E+04		2.95E+04		2.19E+05	

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# NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Metals, Anions, and Miscellaneous (con't)

Analyte	Units	Method Number	Sample Log # 0003012		Sample Log # 0102193		Sample Log # 0106214		Sample Log # 0202251	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1								
Acidity	N	AC7012	3.3E00		3.03E+00		1.74E+00		3.83E00	
Aluminum	μg/L	SW6010B	5.39E+05		1.47E+07		8.43E+06		2.97E+07	
Antimony	μg/L	SW6010B	1.58E+02	U	2.54E+03	В	1.56E+03	В	1.54E+03	В
Arsenic	μg/L	SW6010B	5.17E+02	U	9.08E+02	U	5.8E+02	U	4.65E+02	В
Barium	μg/L	SW6010B	9.9E+01	В	8.39E+03		2.96E+03		2.25E+04	
Beryllium	μg/L	SW6010B	3E00	В	2.02E+02		6.00E+01	В	1.82E+02	
Boron	μg/L	SW6010B							3.71E+05	
Cadmium	μg/L	SW6010B	6.52E+03		5.72E+05		1.47E+05		2.29E+05	
Calcium	μg/L	SW6010B							4.77E+06	
Chloride	μg/L	AC7171							6.86E+05	
Chromium	μg/L	SW6010B	5.44E+03		4.49E+05		8.70E+04		1.67E+05	
Cobalt	μg/L	SW6010B	2.21E+02	В	2.72E+03		1.42E+03	В		
Copper	μg/L	SW6010B	4.57E+02		8.91E+04		1.91E+04			
Fluoride	μg/L	AC7093	4.35E+04	В	2.97E+06	UN	1.00E+06		3.41E+06	
iron	μg/L	SW6010B							1.28E+06	
Lead	μg/L	SW6010B	1.58E+03	В	2.95E+05		8.63E+04		1.00E+05	
Manganese	μg/L	SW6010B	4.85E+03		1.68E+06		3.39E+05		2.97E+05	
Mercury	μg/L	SW7470A	2.24E+04		5.09E+05		1.14E+05		6.28E+06	
Nickel	µg/L	SW6010B	4.20E+03		1.88E+05		5.53E+04		9.49E+04	
Nitrate	μg/L	AC7074							4.42E+08	
Phosphorus	μg/L	SW6010B							2.00E+04	
Potassium	μg/L	SW6010B							2.46E+06	
Selenium	μg/L	SW6010B	8.76E+02	U	5.85E+02	U	9.6E+02	U	3.64E+02	U
Silver	μg/L	SW6010B	3.3E+01	U	3.43E+02	В	4E+02	U	2.47E+03	
Sodium	μg/L	SW6010B							1.50E+07	
Sulfur	μg/L	SW6010B							1.21E+06	
Thallium	μg/L	SW6010B	4.87E+02	U	7.67E+02	U	8E+02	U	4.14E+02	U
Uranium	μg/L	AC7920	5.30E+02		2.11E+05		6.22E+04			
Vanadium	μg/L	SW6010B	3.5E+01	U	1.67E+03	В	5.60E+02	В	8.48E+02	В
Zinc	μg/L	SW6010B	4.62E+02		8.22E+04		3.23E+04		4.34E+04	
Zirconium	μg/L	SW6010B							1.17E+06	
UDS	μg/L	AC7972			1.00E+05		1.29E+06		3.8E+04	
TIC	μg/L	AC8060	4.68E+04	U	1.22E+05	UE	1.19E+05	UE		
TOC	μg/L	SW9060	2.51E+04	В	6.13E+05	В	7.55E+05	В	3.47E+05	

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## NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Analyte	Units	Method Number	Sample L 98120		Sample L 99022		Sample L 99030		Sample L 99031	
			Results	LQF	Results	LQF	Results	LQF	Results	LQI
Semi-Volatile O	rganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	1.90E+02	JBD M	2.5E+01	UM	7.3E+01	М		
2,4-Dinitrotoluene	μg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
2,6-Dinitrotoluene	μg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
4-Nitrophenol	μg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
4,6-Dinitro-2- nethylphenol	μg/L	SW8270C	2E+01	UM	3.1E+01	М	2.5E+01	U		
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	1.90E+02	BZD	3.4E+01		5.40E+02	D		
Butylbenzyl phthalate	μg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
Diethylphthalate	μg/L	SW8270C	4.6E+01		2.5E+01	UM	2.5E+01	UM		
Di-n-octyl phthalate	µg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
Nitrobenzene	μg/L	SW8270C	2E+01	U	4.1E+01	М	2.5E+01	UM		
n- Nitrosodimethylamine	μg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
Pyridine	μg/L	SW8270C	2E+01	U	2.5E+01	U	2.5E+01	U		
ri-n-butyl phosphate	μg/L	SW8270C	2E+01	υ	2.5E+01	UM	2.5E+01	UM		
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/ <b>L</b>	SW8260A	1E+01	U	2E00	UН	2E00	U	3E00	U I M
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	3E00	UH
2-Butanone	μg/L	SW8260A	1.20E+01		3E00	UН	3E00	U	8E00	UH
2-Hexanone	μg/L	SW8260A	3E00	J M	3E00	U H M	3E00	UM	2.3E+01	U ł M
1-Methyl-2-pentanone	μ <b>g/l</b> L	SW8260A	1E+01	U	3E00	U H M	3E00	UM	1.2E+01	U F M
Acetone	μg/L	SW8260A	1E+01	U	2E00	JH	2E00	U	4E+00	UH
3enzene	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	3E00	U F M
3romodichloromethane	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	3E00	UH
Bromoform	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	1.4E+01	UH
Bromomethane	μg/L	SW8260A	1E+01	U	6E00	JH	3E00	J	1.2E+02	B H M
Carbon disulfide	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	4E00	UH
Carbon tetrachloride	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	υ	7E00	UH
Chloroform	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	υ	3E00	Uŀ
Chloromethane	μg/L	SW8260A	1E+01	U	2E00	JHM	2E00	UM	1.7E+01	U F M
Dibromochloromethane	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	6E00	UH
Dichloromethane	μg/L	SW8260A	1E+01	UM	2E00	U H M	2E00	UM	8.6E+01	нк
Styrene	μg/L	SW8260A	1E+01	UM	2E00	U H M	2E00	UM	1.4E+01	U F M
Trichlorofluoromethane	μg/L	SW8260A	1E+01	U	2E00	UН	2E00	U	4E00	UH

μg/L

21 / 3.79E+03

21 / 6.23E+02

17 / 5.78E+02

21 / 3.74E+03

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## NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Analyte	Units	Method Number	Sample L 99032		Sample L 99040	-	Sample L 99041	-	Sample L 99062	
			Results	LQF	Results	LQF	Results	LQF	Results	LQ
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	2.7E+02	M D	2.9E+01	М	2.5E+01	U M	2.5E+01	UN
2,4-Dinitrotoluene	μg/L	SW8270C	5E+01	U D	2.5E+01	U	2.5E+01	U	2.5E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	5E+01	U D	2.5E+01	U	2.5E+01	U	2.5E+01	U
4-Nitrophenol	μg/L	SW8270C	5E+01	U D	2.5E+01	U	2.5E+01	U	2.5E+01	U
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	5E+01	U M D	2.5E+01	UM	2.5E+01	UM	2.5E+01	UN
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	5E+01	UD	4.5E+01	В	6.8E+01	В	3.6E+01	
Butylbenzyl phthalate	μg/L	SW8270C	5E+01	U D	2.5E+01	U	2.5E+01	U	2.5E+01	U
Diethylphthalate	μg/ <b>L</b>	SW8270C	5E+01	U M D	2.5E+01	UM	2.5E+01	UM	2.5E+01	UN
Di-n-octyl phthalate	μg/L	SW8270C	5E+01	U D	2.5E+01	UΖ	2.5E+01	U	2.5E+01	U
Nitrobenzene	μg/L	SW8270C	5E+01	D D	2.5E+01	UM	2.5E+01	UM	2.5E+01	UN
n- Nitrosodimethylamine	μg/L	SW8270C	5E+01	U M D	2.5E+01	U M	2.5E+01	U	2.5E+01	U
Pyridine	μ <b>g/L</b>	SW8270C	5E+01	U D	2.5E+01	U	2.5E+01	U	2.5E+01	U
Fri-n-butyl phosphate	μg/L	SW8270C	5E+01	U M D	2.5E+01	UM	2.5E+01	UM	2.5E+01	U
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	3E00	U M	3E00	UM	3E00	UM	2E00	U
1,1,1-Trichloroethane	µg/L	SW8260A	3E00	U	3E00	U	3E00	U	2E00	U
2-Butanone	μg/L	SW8260A	8E00	U	8E00	U	8E00	U	3E00	U
2-Hexanone	μ <b>g/L</b>	SW8260A	2.3E+01	UM	2.3E+01	UM	2.3E+01	UM	3E00	U
4-Methyl-2-pentanone	µg/L	SW8260A	1.2E+01	UM	1.2E+01	UM	1.2E+01	U M	3E00	U
Acetone	μg/L	SW8260A	4E00	U	6E00	J	4E00	JB	9E00	JВ
Benzene	µg/L	SW8260A	3E00	υ <b>м</b>	3E00	UM	3E00	UM	2E00	UN
Bromodichloromethane	µg/L	SW8260A	3E00	U	3E00	υ	3E00	U	2E00	U
Bromoform	µg/L	SW8260A	1.4E+01	U	1.4E+01	υ	1.4E+01	U	2E00	U
Bromomethane	μg/L	SW8260A	5.9E+01	вм	9.5E+01	М	4.7E+01	ВМ	7E00	JM
Carbon disulfide	μg/L	SW8260A	4E00	U	4E00	U	4E00	U	2E00	U
Carbon tetrachloride	μg/L	SW8260A	7E00	U	7E00	U	7E00	U	2E00	U
Chloroform	µg/L	SW8260A	3E00	U	3E00	U	3E00	U	2E00	u
Chloromethane	μg/L	SW8260A	1.7E+01	UM	8.9E+01	М	1.7E+01	UM	2E00	U
Dibromochloromethane	μg/L	SW8260A	6E00	U	6E00	U	6E00	U	2E00	U
Dichloromethane	μg/L	SW8260A	6E00	UM	6E00	UM	2.1E+01	М	2E00	U
Styrene	μg/L	SW8260A	1.4E+01	UM	1.4E+01	UM	1.4E+01	U M	2E00	UΙ
Trichlorofluoromethane	μg/L	SW8260A	4E00	υ	4E00	U	<b>4</b> E00	U	2E00	U

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## NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Analyte	Units	Method Number	Sample Le 990907		Sample Lo 991019		Sample L 991107		Sample L 000112	
			Results	LQF	Results	LQF	Results	LQF	Results	LQ
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	8E+01	М	4E+01	υM	9.9E+01	М	4E+01	D D
2,4-Dinitrotoluene	µg/L	SW8270C	2.5E+01	U	2E+01	ť	2E+01	U	4E+01	UE
2,6-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2E+01	U	2E+01	U	4E+01	υŒ
4-Nitrophenol	μg/L	SW8270C	2.5E+01	U	2E+01	U	1E+01	U	4E+01	Ų
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	2.5E+01	U	4E+01	U M	4E+01	U M	4E+01	U N
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	2.5E+01	U	2E+01	U	2E+01	บ	4E+01	U
Butylbenzyl phthalate	μg/L	SW8270C	2.5E+01	υ	2E+01	U	2E+01	U	4E+01	U
Diethylphthalate	μg/L	SW8270C	2.5E+01	UM	2E+01	UM	2E+01	U M	4E+01	U
Di-n-octyl phthalate	μg/L	SW8270C	2.5E+01	U	2E+01	U	2E+01	U	4E+01	UE
Nitrobenzene	μg/L	SW8270C	2.5E+01	UM	4E+01	U	2E+01	U	4E+01	UE
n- Nitrosodimethylamine	μg/L	SW8270C	2.5E+01	U	4E+01	U	4E+01	บ	4E+01	U
Pyridine	µg/L	SW8270C	2.5E+01	υ	2E+01	U	2E+01	U	4E+01	U
Tri-n-butyl phosphate	μg/L	SW8270C	2.5E+01	UM	2E+01	UМ	2E+01	υм	4E+01	U
Volatile Organic			2.32.101	0 101	22.01		22.101	0 141	42.01	
1,1-Dichloroethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E00	υ
1,1,1-Trichloroethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E00	U
2-Butanone	μg/L	SW8260A	2E00	U	2E00	U	1E+01	U	2E00	UN
2-Hexanone	μg/L	SW8260A	2E00	U	2E00	U	1E+01	U	2E00	U
					2E00	U	1E+01	U	2E00	UI
4-Methyl-2-pentanone	μg/L 	SW8260A	2E00	U						
Acetone	µg/L	SW8260A	4E00	JBM	2E00	U	3E00	J B	4E00	J N
Benzene	µg/L	SW8260A	1E00	UM	1E00	UM	3E00	JBM	1E00	UN
Bromodichloromethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E00	U
Bromoform	µg/L	SW8260A	2E00	UM	2E00	U	1E+01	U	2E00	U
Bromomethane	μg/L	SW8260A	2E00	U	2E00	U	1E+01	U	2E00	U
Carbon disulfide	µg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E00	U
Carbon tetrachloride	μg/ <b>L</b>	SW8260A	2E00	U	2E00	U	1E+01	U	2E00	U
Chloroform	µg/Ł	SW8260A	1E00	υ	1E00	U	1E+01	U	1E00	U
Chloromethane	μg/L	SW8260A	2E00	U	2E00	U	1E+01	U	2E00	UI
Dibromochloromethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E00	U
Dichloromethane	µg/L	SW8260A	1E00	U	1E00	UM	1E+01	UM	1E00	บเ
Styrene	μg/L	SW8260A	1E00	UM	1E00	UΜ	1E+01	UM	1E00	UI
Trichlorofluoromethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E00	U
Tentatively Ident	ified Or	ganic Comp	ounds							
no. Ided / total conc.	μg/L		15 / 6.48E+02	SVOCs only	10 / 8.88E+02		11 / 3.27E+02		1 / 1.80E+04	

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## NWCF Fluoride Hot Sump Tank - NCC-119 (con't.)

Analyte	Units	Method Number	Sample L 00030		Sample I 01021		Sample Lo 010621		Sample L 02022	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
Semi-Volatile Or	rganic C	ompounds								
2,4-Dinitrophenol	µg/L	SW8270C	9E00	J	2E+01	υM	4.4E+01	М	1.4E+01	UM
2,4-Dinitrotoluene	μg/L	SW8270C	2E+01	U	2E+01	U	2E+01	U	8E00	U
2,6-Dinitrotoluene	μg/L	SW8270C	2E+01	U	1.2E+01	J	2E+01	U	1.1E+01	U
4-Nitrophenol	μg/L	SW8270C	2E+01	U	2E+01	U	2E+01	UM	3E+01	UΧ
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	2E+01	U	2E+01	UM	2E+01	UM	1.3E+01	UM
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	6E00	J	2E+01	U	1.4E+02	U	9E00	UM
Butylbenzyl phthalate	μg/L	SW8270C	2E+01	U	2E+01	U	2E+01	U	9E00	υм
Diethylphthalate	μg/L	SW8270C	8E00	J M	2E+01	UM	2E+01	UM	1.2E+01	UM
Di-n-octyl phthalate	µg/L	SW8270C	2E+01	U	2E+01	U	2E+01	U	6E00	U
Nitrobenzene	μg/L	SW8270C	2E+01	UM	2E+01	U	2E+01	U	7E00	υм
n- Nitrosodimethylamine	μg/L	SW8270C	2E+01	UM	1.4E+02		3.6E+01		2.4E+01	υx
Pyridine	μg/L	SW8270C	2E+01	U	2E+01	U	2E+01	U	1.5E+01	U
Tri-n-butyl phosphate	μg/L	SW8270C	2E+01	UM	2E+01	U			1.1E+01	ВЈМ
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
2-Butanone	μg/L	SW8260A	1E+01	UM	1E+01	U	1E+01	U	1E+01	U
2-Hexanone	μg/L	SW8260A	1E+01	υM	2E+01	U M Z	2E+01	υz	1E+01	U
4-Methyl-2-pentanone	μg/L	SW8260A	1E+01	U	2E+01	UΖ	2E+01	υz	1E+01	U
Acetone	μg/L	SW8260A	7E00	вјм	2E+01	UZY	2E+01	υZ	7E00	JB
Benzene	μg/L	SW8260A	1E+01	UM	1E+01	U	1E+01	U	1E+01	U M
Bromodichloromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Bromoform	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Bromomethane	μg/L	SW8260A	1E+01	U	1.8E+01		3.7E+01	вм	2.6E+01	
Carbon disulfide	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Carbon tetrachloride	μg/L	SW8260A	1E+01	U	1E+01	UM	1E+01	U	1E+01	UM
Chloroform	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Chloromethane	μg/L	SW8260A	1E+01	U	6E00	J	1E+01	U	1.7E+01	M
Dibromochloromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Dichloromethane	μg/L	SW8260A	4.8E+01	М	2E+01	UZY	2E+01	UΖ	1E+01	U
Styrene	μ <b>g/L</b>	SW8260A	1E+01	UM	1E+01	U	2E+01	UΖ	1E+01	UM
Trichlorofluoromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Tentatively Iden	tified Or	ganic Comp	ounds							
no. Ided / total conc.	μg/L		4 / 4.91E+02		9 / 2.94E+02	SVOCs only	21 / 5.22E+02		12 / 4.94E+02	



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## **ENGINEERING DESIGN FILE**

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## NWCF Non-Fluoride Hot Sump Tank - NCC-122

Metals, Anions, and Miscellaneous

Analyte	Units	Method Number	Sample L 990205		Sample Li 990418	•	Sample L 000328	-	Sample L 010212	_
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1	0.35		0.63	В				
Acidity	N	AC7012					3.3E-01	В	4.1E-01	
Aluminum	μg/L	SW6010B	2.06E+04		6.72E+05		1.88E+05		2.01E+04	
Antimony	μg/L	SW6010B	4.52E+02	U	2.20E+02	U	1.32E+02	U	9E00	В
Arsenic	μg/L	SW6010B	5.04E+02	U	2.37E+02	U	8.5E+01	U	4.5E00	U
Barium	μg/L	SW6010B	1.10E+02	В	1.17E+02	В	4.13E+02		1.54E+01	
Beryllium	μg/L	SW6010B	2.0E00	В	3.0E00	В	1.7E00		3E-01	В
Boron	µg/L	SW6010B								
Cadmium	μg/L	SW6010B	2.76E+02	В	2.11E+03	N	4.55E+02		7.88 <b>E+</b> 02	
Calcium	μg/L	SW6010B								
Chloride	µg/L	AC7171								
Chromium	µg/L	SW6010B	2.24E+03		4.29E+03	N	1.27E+03		6.57E+02	
Cobalt	µg/L	SW6010B	7.4E+01	U	2.4E+01	В	1.67E+01		5.7E00	В
Copper	μg/L	SW6010B	5.24E+02	В	4.83E+02		8.60E+02		1.39E+02	
Fluoride	μg/L	AC7093	2.56E+04	UE	3.31E+04	В	1.85E+04	U	1.14E+05	UN
Iron	µg/L	SW6010B								
Lead	μg/L	SW6010B	3.72E+02	U	1.92E+03	В	8.60E+02		4.19E+02	
Manganese	μg/L	SW6010B	3.64E+03		7.32E+03		2.33E+03		2.17E+03	
Mercury	μg/L	SW7470A	1.91E+04		6.34E+05		1.90E+03		7.44E+03	
Nickel	μg/L	SW6010B	1.81E+03		1.60E+03		9.07E+02		3.01E+02	
Nitrate	μg/L	AC7074								
Phosphorus	µg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	3.32E+02	U	2.67E+02	U	7.8 <b>E+01</b>	U	2.9E00	U
Silver	μg/L	SW6010B	1.28E+02	U	6.2E+01	U	2.5E+01	U	1.7E00	U
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	4.68E+02	U	2.84E+02	U	1.18E+02	U	3.8E00	U
Uranium	μg/L	AC7920	1.89E+02	U	3.17E+02		1.29E+02		3.42E+02	
Vanadium	μg/L	SW6010B	1.64E+02	U	3.6E+01	U	4.5E+01	U	3.4E+00	В
Zinc	μg/L	SW6010B	2.57E+03		2.88E+03		5.78E+03		2.17E+02	
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972					0.00E00		2E+05	
TIC	μg/L	AC8060	5.83E+04	U	5.82E+04	U	1.40E+05	UE	2.38E+04	U E
тос	μg/L	SW9060	4.95E+04		1.18E+05	В	8.20E+04	В	7.92E+04	В

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## NWCF Non-Fluoride Hot Sump Tank - NCC-122 (con't.)

## Metals, Anions, and Miscellaneous (con't.)

Analyte	Units	Method Number	Sample Le 010622	•	Sample Lo 020214					
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
рН		EPA150.1								
Acidity	N	AC7012	5.1E-01		7.19E-01					
Aluminum	μg/L	SW6010B	1.7E+03		8.63E+02					
Antimony	µg/L	SW6010B	4.7E+01	U	3.7E+01	U				
Arsenic	μg/L	SW6010B	2.9E+01	U	3.3E+01	U				
Barium	μg/L	SW6010B	9E00	В	1.0E+01	В				
Beryllium	μg/L	SW6010B	1E00	U	1E00	U				
Boron	μg/L	SW6010B			3.56E+03					
Cadmium	μg/L	SW6010B	4E00	U	6.0E00	В				
Calcium	μg/L	SW6010B			9.84E+02	Ε				
Chloride	μg/L	AC7171			3.24E+04	U				
Chromium	μg/L	SW6010B	2.7E+01	В	5.5E+01	В				
Cobalt	μg/L	SW6010B	1E+01	U						
Copper	μg/L	SW6010B	1.4E+01	U						
Fluoride	μg/L	AC7093	7.02E+04	U	2.35E+04	U				
Iron	μg/L	SW6010B			2.32E+03					
Lead	μg/L	SW6010B	6.3E+01	U	4.4E+01	U				
Manganese	μg/L	SW6010B	1.3E+01	В	3.2E+01					
Mercury	μg/L	SW7470A	3.95E+03		1.66E+04					
Nickel	μg/L	SW6010B	3.2E+01	В	6.0E+01	В				
Nitrate	μg/L	AC7074			4.04E+07					
Phosphorus	μg/L	SW6010B			8.1E+01	В				
Potassium	μg/L	SW6010B			2.62E+02	U				
Selenium	μg/L	SW6010B	4.8E+01	U	3.6E+01	U				
Silver	μg/L	SW6010B	2E+01	U	2.0E+01	U				
Sodium	μg/L	SW6010B			4.12E+03					
Sulfur	μg/L	SW6010B			2.62E+02	В				
Thallium	μg/L	SW6010B	4E+01	U	4.1E+01	U				
Uranium	mg/L	AC7920	3.2E+02	U						
Vanadium	μg/L	SW6010B	1E+01	U	1.4E+01	U				
Zinc	μg/L	SW6010B	5.3E+01		4.1E+01					
Zirconium	μg/L	SW6010B			6.0E+01	В				
UDS	μg/L	AC7972	5.0E+03	U	1E+05					
TIC	μg/L	AC8060	2.38E+04	UE						
тос	μg/L	SW9060	1.48E+05	В	2.94E+04					

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## NWCF Non-Fluoride Hot Sump Tank - NCC-122 (con't.)

Analyte	Units	Method Number	Sample L 990205		Sample L 99041		Sample L 000328		Sample L 010212	
			Results	LQF	Results	LQF	Results	LQF	Results	LQ
Semi-Volatile Orç	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C			2.5E+01	UM	7.8E+01	М	1.2E+02	М
2,4-Dinitrotoluene	μg/L	SW8270C			2.5E+01	U	2E+01	U	7E00	J
2,6-Dinitrotoluene	µg/L	SW8270C			2.5E+01	U	2E+01	U	2E+01	U
4-Nitrophenol	μg/L	SW8270C			2.5E+01	U	1.8E+01	J M	2E+01	U
4,6-Dinitro-2- methylphenol	μg/L	SW8270C			2.5E+01	UM	2E+01	UM	2E+01	UN
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C			2.5E+01	U	4E+01		5.2E+01	
Butylbenzyl phthalate	μg/L	SW8270C			2.5E+01	U	2E+01	U	2E+01	U
Diethylphthalate	μg/L	SW8270C			2.5E+01	UM	1E+01	J	2E+01	UM
Di-n-octyl phthalate	μg/L	SW8270C			2.5E+01	U	2E+01	U	5.8E+01	
Nitrobenzene	μg/L	SW8270C			2.5E+01	UM	2E+01	U	2E+01	J
n- Nitrosodimethylamine	μg/L	SW8270C			2.5E+01	UM	2E+01	U	3.6E+02	D
Pyridine	μg/L	SW8270C			2.5E+01	U	2E+01	U	2E+01	U
Tri-n-butyl phosphate	μg/L	SW8270C			2.5E+01	UM	2E+01	UM	4.8E+01	
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A			3E00	UM	1E+01	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A			3E00	U	1E+01	U	1E+01	U
2-Butanone	μ <b>g/L</b>	SW8260A			8E00	U	1E+01	U	1E+01	U
2-Hexanone	μg/L	SW8260A			2.3E+01	UM	1E+01	U	1E+01	U
4-Methyl-2-pentanone	μg/L	SW8260A			1.2E+01	UM	1E+01	U	1E+01	U
Acetone	μg/L	SW8260A			4E00	U	1.8E+01	вм	1.1E+02	E
Benzene	μg/L	SW8260A			3E00	U M	1E+01	UM	1E+01	U
Bromodichloromethane	μ <b>g/L</b>	SW8260A			3E00	U	1E+01	U	1E+01	U
Bromoform	μg/L	SW8260A			1.4E+01	U	1E+01	U	1E+01	U
Bromomethane	μg/L	SW8260A			7E00	UM	1E+01	U M	1.9E+01	
Carbon disulfide	μg/L	SW8260A			4E00	U	1E+01	U	1E+01	U
Carbon tetrachloride	μg/L	SW8260A			7E00	U	1E+01	U	1E+01	U
Chloroform	μg/ <b>L</b>	SW8260A			3E00	U	1E+01	U	1E+01	U
Chloromethane	μ <b>g/L</b>	SW8260A			1.7E+01	UМ	1E+01	U	3.7E+01	M
Dibromochloromethane	µg/L	SW8260A			6E00	U	1E+01	U	1E+01	υ
Dichloromethane	μg/L	SW8260A			6E00	UM	1E+01	UM	1E+01	U
Styrene	μg/L	SW8260A			1.4E+01	UM	1E+01	UM	1E+01	U
Trichlorofluoromethane	μg/L	SW8260A			4E00	U	1E+01	U	1E+01	U
Tentatively Ident			ounds							
•		• '				SVOCs				

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## NWCF Non-Fluoride Hot Sump Tank - NCC-122 (con't.)

Volatile Organic Compounds and Semi-volatile Organic Compounds (con't.)

Analyte	Units	Method Number	Sample L 01062		Sample L 020214					
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
Semi-Volatile Org	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	1.1E+02	М	1.3E+01	U M				
2,4-Dinitrotoluene	μg/L	SW8270C	2E+01	U	8E00	U				
2,6-Dinitrotoluene	μg/L	SW8270C	2E+01	U	1.1E+01	U				
4-Nitrophenol	μg/L	SW8270C	2E+01	UM	3E+01	υx				
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	2E+01	UM	1.3E+01	UM				
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	2E+01	U	9E00	UM				
Butylbenzyl phthalate	μg/L	SW8270C	2E+01	U	9E00	υM				
Diethylphthalate	μg/L	SW8270C	2E+01	UM	1.2E+01	υM				
Di-n-octyl phthalate	μg/L	SW8270C	2E+01	U	6E00	U				
Nitrobenzene	μg/L	SW8270C	2E+01	U	7E00	UM				
n- Nitrosodimethylamine	μ <b>g/L</b>	SW8270C	4.2E+01		2.4E+01	UX				
Pyridine	μ <b>g/L</b>	SW8270C	2E+01	υ	1.5E+01	U				
Tri-n-butyl phosphate	μg/L	SW8270C			4.2E+01	ВМ				
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U				
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U				
2-Butanone	µg/L	SW8260A	1E+01	U	1E+01	U				
2-Hexanone	μ <b>g/L</b>	SW8260A	2E+01	UΖ	1E+01	U				
4-Methyl-2-pentanone	μg/L	SW8260A	2E+01	UΖ	1E+01	U				
Acetone	μg/L	SW8260A	3.2E+01	Y	5.1E+01	В				
Benzene	μg/L	SW8260A	1E+01	U	1E+01	UM				
Bromodichloromethane	μg/L	SW8260A	1E+01	U	1E+01	U				
Bromoform	μg/L	SW8260A	1E+01	U	1E+01	U				
Bromomethane	µg/L	SW8260A	1.6E+02	E B M	4E+01					
Carbon disulfide	μg/L	SW8260A	1E+01	U	1E+01	U				
Carbon tetrachloride	µg/L	SW8260A	1E+01	U	1E+01	UM				
Chloroform -	μg/L	SW8260A	1E+01	U	1E+01	U				
Chloromethane	μ <b>g/L</b>	SW8260A	1E+01	U	7E+01	М				
Dibromochloromethane	µg/L	SW8260A	1E+01	U	1E+01	U				
Dichloromethane	μg/L	SW8260A	2E+01	υz	1E+01	U				
Styrene	μg/L	SW8260A	1E+01	U	1E+01	U M				
Trichlorofluoromethane	μg/L	SW8260A	1E+01	U	1E+01	U				

7 / 1.65E+02

no. Ided / total conc. μg/L

9 / 1.61E+02



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## **ENGINEERING DESIGN FILE**

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## NWCF Decontamination Hold Tanks - NCD-123 & NCD-129

Metals, Anions, and Miscellaneous

Analyte	Units	Method Number	Sample Lo 990210		Sample Lo 990402		Sample Le 990617		Sample L 010417	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1	0.97		0.34	U				
Acidity	N	AC7012					5.7E-01	В	7.79E-01	
Aluminum	μg/L	SW6010B	9.14E+04		7.39E+04		5.25E+04		7.14E+03	
Antimony	μg/L	SW6010B	3.12E+02	В	2.20E+02	U	4.57E+02		2.06E+03	
Arsenic	μg/L	SW6010B	2.52E+02	U	2.37E+02	U	3.9E+01	U	1.45E+01	U
Barium	μg/L	SW6010B	9.5E+01	В	1.07E+02	В	3.39E+02		2.26E+03	
Beryllium	μg/L	SW6010B	2.0E00	В	2E00	U	1E00	U	5E-01	U
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	6.94E+02		3.81E+02		1.35E+02		4.5E+01	
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171								
Chromium	μg/L	SW6010B	4.21E+03		4.88E+03		2.13E+03		5.78E+02	
Cobalt	μg/L	SW6010B			2.2E+01	U	2.3E+01	В	5E00	U
Copper	μg/L	SW6010B			1.32E+03		3.10E+03		1.6E+02	
Fluoride	μg/L	AC7093	2.56E+04	UE	2.57E+04	U	1.28E+04	U	7.02E+04	U
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	4.11E+02	В	2.16E+03		1.92E+03		1.16E+03	
Manganese	μg/L	SW6010B			4.38E+04		1.24E+04		5.36E+02	
Mercury	μg/L	SW7470A	3.63E+02		1.03E+03		2.93E+03	E	2.12E+02	
Nickel	μg/L	SW6010B	8.58E+02		2.21E+03		1.23E+03		3.47E+02	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	1.66E+02	U	2.67E+02	υ	4.3E+01	U	2.4E+01	U
Silver	μg/L	SW6010B	6.4E+01	U	1.08E+01	В	7.7E+01	В	1E+01	U
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	2.34E+02	U	2.84E+02	U	3.4E+01	U	2E+01	U
Uranium	μg/L	AC7920	1.90E+02	U					3.2E+02	U
Vanadium	μg/L	SW6010B	8.2E+01	U	5.0E+01	В	3.5E+01	В	2.05E+01	В
Zinc	μg/L	SW6010B	6.98E+03		2.13E+04		1.16E+04		3.1E+03	
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972							2.5E+04	
TIC	μg/L	AC8060	9.32E+04	U					5.95E+04	UΕ
тос	μg/L	SW9060	1.24E+06						1.66E+05	

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## NWCF Decontamination Hold Tanks - NCD-123 & NCD-129 (con't.)

## Metals, Anions, and Miscellaneous (con't.)

Analyte	Units	Method Number	Sample Lo 010702				
			Results	LQF	 	 	
рН		EPA150.1					
Acidity	N	AC7012	7.2E-01				
Aluminum	μg/L	SW6010B	2.14E+04				
Antimony	μg/L	SW6010B	3.24E+02				
Arsenic	μg/L	SW6010B	1.45E+01	U			
Barium	μg/L	SW6010B	3.03E+03				
Beryllium	μg/L	SW6010B	5E-01	В			
Boron	μg/L	SW6010B					
Cadmium	μg/L	SW6010B	1.85E+02				
Calcium	μg/L	SW6010B					
Chloride	μg/L	AC7171					
Chromium	μg/Ľ	SW6010B	4E+03				
Cobalt	μg/L	SW6010B	3.85E+01	В			
Copper	μg/L	SW6010B	4E+02				
Fluoride	μg/L	AC7093	7.02E+04	U			
Iron	μg/L	SW6010B					
Lead	μg/L	SW6010B	2.02E+02	В			
Manganese	μg/L	SW6010B	8.06E+02				
Mercury	μg/L	SW7470A	1.21E+03				
Nickel	µg/L	SW6010B	2.93E+03				
Nitrate	μg/L	AC7074					
Phosphorus	μg/L	SW6010B					
Potassium	μg/L	SW6010B					
Selenium	μg/L	SW6010B	2.4E+01	U			
Silver	µg/L	SW6010B	2.22E+02				
Sodium	μg/L	SW6010B					
Sulfur	μg/L	SW6010B					
Thallium	μg/L	SW6010B	2E+01	U			
Uranium	μg/L	AC7920	3.2E+02	U			
Vanadium	μg/L	SW6010B	2.7E+01	В			
Zinc	μg/L	SW6010B	2.27E+03				
Zirconium	μg/L	SW6010B					
UDS	μg/L	AC7972	5.0E+03	U			
TIC	μg/L	AC8060	2.38E+04	UE			
TOC	μg/L	SW9060	1.98E+04	UE		 	

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## NWCF Decontamination Hold Tanks - NCD-123 & NCD-129 (con't.)

Analyte	Units	Method Number	Sample L 990210		Sample L 990402		Sample L 99061		Sample I 01041	
			Results	LQF	Results	LQF	Results	LQF	Results	LQ
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	µg/L	SW8270C	7.6E+01	М	2.5E+01	UM	2.6E+01	М	2E+01	UN
2,4-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
4-Nitrophenol	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	2.5E+01	U	2.5E+01	UM	2E+01	UM	2E+01	U
Bis-(2-ethylhexyl) ohthalate	μg/L	SW8270C	4.7E+01		2.5E+01	U	2E+01	U	2E+01	U
Butylbenzyl phthalate	µg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	u
Diethylphthalate	μg/L	SW8270C	2.5E+01	υм	2.5E+01	υM	2E+01	U	2E+01	U
Di-n-octyl phthalate	μg/L	SW8270C	2.5E+01	U	2.5E+01	υz	2E+01	U	2E+01	U
Nitrobenzene	μg/L	SW8270C	2.5E+01	U	2.5E+01	υ	2E+01	U M	2E+01	U
n- Nitrosodimethylamine	μg/L	SW8270C	2.5E+02	UD	2.5E+01	UM	2E+01	U	2E+01	u
Pyridine	μg/L	SW8270C	2.5E+02	UD	2.5E+01	U	2E+01	U	2E+01	L
Fri-n-butyl phosphate	μg/L	SW8270C	2.5E+01	UM	2.5E+01	υм	2E+01	U	2E+01	U
Volatile Organic		unds								
1,1-Dichloroethane	µg/L	SW8260A	5E+01	U D H	1E+01	U M	1E+01	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	5E+01	U D H	1E+01	U	1E+01	U	1E+01	ι
2-Butanone	μg/L	SW8260A	5E+01	U D H	1E+01	υ	1E+01	U	1E+01	u
2-Hexanone	μg/L	SW8260A	5E+01	U D H	5E+01	UM	1E+01	U	1E+01	u
4-Methyl-2-pentanone	μg/L	SW8260A	5E+01	U D H	5E+01	U M Z	1E+01	U	1E+01	u
Acetone	µg/L	SW8260A	1.3E+01	J D H	1E+01	- U	8E00	JBM	1.4E+01	
Benzene	μg/L	SW8260A	5E+01	U D H M	1E+01	UM	1E+01	UM	1E+01	u
				UD				U		u
Bromodichloromethane	μg/L	SW8260A	5E+01	H U D	1E+01	U	1E+01		1E+01	
Bromoform	µg/L	SW8260A	5E+01	Н U D	2E+01	υz	1E+01	U	1E+01	U
Bromomethane	μg/L	SW8260A	5E+01	H M U D	1E+01	UM	1E+01	UM	1E+01	u
Carbon disulfide	μ <b>g/L</b>	SW8260A	5E+01	Н	1E+01	U	1E+01	U	1E+01	U
Carbon tetrachloride	μg/L	SW8260A	5E+01	H	1E+01	U	1E+01	U	1E+01	ι
Chloroform	μg/L	SW8260A	5E+01	H H	1E+01	U	1E+01	U	1E+01	L
Chloromethane	μg/L	SW8260A	5E+01	U D H	2E+01	U M Z	1E+01	U	1E+01	U
Dibromochloromethane	μg/L	SW8260A	5E+01	U D H	1E+01	U	1E+01	U	1E+01	L
Dichloromethane	μg/L	SW8260A	5E+01	H H	1E+01	υм	1E+01	U	1E+01	, u
Styrene	μg/L	SW8260A	5E+01	U D H <b>M</b>	2E+01	UM Z	1E+01	UM	1E+01	u
Trichlorofluoromethane	μg/L	SW8260A	5E+01	U D H	1E+01	U	1E+01	U	1E+01	U
Tentatively Ident	ified Or	ganic Comp	oounds							
no. Ided / total conc.	μg/L		20 / 1.03E+04	SVOCs only	6 / 2.21E+02		12 / 3.99E+02	SVOCs only	5 / 3.05E+02	SVO(

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## NWCF Decontamination Hold Tanks - NCD-123 & NCD-129 (con't.)

Analyte	Units	Method Number	Sample L 01070							
			Results	LQF	Results	LQF	Results	LQF	Results	LQI
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	µg/L	SW8270C	2E+01	U H M						
2,4-Dinitrotoluene	μg/L	SW8270C	2E+01	UН						
2,6-Dinitrotoluene	μg/L	SW8270C	2E+01	UН						
4-Nitrophenol	μg/L	SW8270C	2E+01	<b>м</b>						
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	2E+01	UH M						
Bis-(2-ethylhexyl)										
phthalate	µg/L	SW8270C	2E+01	UН						
Butylbenzyl phthalate	μg/L	SW8270C	2E+01	UН						
Diethylphthalate	μg/L	SW8270C	2E+01	U H M						
Di-n-octyl phthalate	μg/L	SW8270C	2E+01	υн						
Nitrobenzene	μg/L	SW8270C	2E+01	UН						
n- Nitrosodimethylamine	μg/L	SW8270C	2E+01	UН						
Pyridine	μg/L	SW8270C	2E+01	UН						
Tri-n-butyl phosphate	μg/L	SW8270C								
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	1E+01	U						
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U						
2-Butanone	μg/L	SW8260A	1E+01	υ						
2-Hexanone	μg/L	SW8260A	1E+01	U						
4-Methyl-2-pentanone	μg/L	SW8260A	1E+01	U						
Acetone	μg/L	SW8260A	1E+01	U						
Benzene	μg/L	SW8260A	1E+01	U						
Bromodichloromethane	μg/L	SW8260A	1E+01	U						
Bromoform	μg/L	SW8260A	1E+01	U						
Bromomethane	μg/ <b>L</b>	SW8260A	1E+01	U						
Carbon disulfide	μg/L	SW8260A	1E+01	U						
Carbon tetrachloride	μg/L	SW8260A	1E+01	U						
Chloroform	μg/L	SW8260A	1E+01	U						
Chloromethane	μg/L	SW8260A	1E+01	UM						
Dibromochloromethane	μg/L	SW8260A	1E+01	U						
Dichloromethane	μg/L	SW8260A	6E00	JBM						
Styrene	μg/L	SW8260A	1E+01	U						
Trichlorofluoromethane	μg/L	SW8260A	1E+01	U		-				
Tentatively Iden	tified Or	ganic Comp	ounds							
				SVOCs						



Log Search

Date of Search: 2003-06-16 13:42:09.522 Run by: JEFF LONG \* Search Criteria: Start Log....:980624 1 End Log....:980624 1 Log Approval .: ALL Logs Result Type..: All Entries Lab/Group....:ALL Groups Name Column..:Lab Sample ID Total # Logs Found...: 1 Total # Results Found: 9 \* Request Name Log Type Charge Num Log Approval Info Log # I L d Lab Meth a Anax ID # b lyst Analyte ARL Result 56161B201 011010 16:29 KIMBERLY A WHITEH 980624-1 WL-133 PLANT 57171 7BGP CHLORIDE A.. 5.1E+01 +- 1.6E+01 ug/mL 1 8BP53 57171 7BGP CHLORIDE A.. 5.1E+01 +- 1.6E+01 ug/mL
57093 7BGP FLUORIDE A.. 8.49E+01 +- 6.2E+00 ug/mL
97074 7BCS NITRATE A.. 6.58E-01 +- 8.0E-02 Molar
97168 7BCS SULFATE A.. 2.61E+02 +- 2.2E+01 ug/ml
87100 7BCS ALUMINUM A.. 3.61E-02 +- 1.2E-03 MOLAR
57015 7BGP ACID A.. 4.0E-01 +- 1.8E-01 Normal Acid
47981 7BGP SPGR A.. 1.02375E+00 +- 6.6E-05 @ 25/4
17920 7BCS URANIUM A.. 3.42E-03 +- 3.3E-04 G/L
17972 7BGP UDS A.. 43.524 G/L 8BP53 2 8BP53 3 8BP53 8BP53 6 8BP53 8BP53 7 8 8BP53 9 8BP53

\*\*\*\*\*\*\* END \*\*\*\*\*\*

Log Search

Date of Search: 2003-06-16 13:44:19.946 Run by: JEFF LONG \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

Search Criteria:

Start Log....:960928 1 End Log....:960928 1 Log Approval.:ALL Logs Result Type..:All Entries Lab/Group....:ALL Groups Name Column..:Lab Sample ID

Total # Logs Found...: 1 Total # Results Found: 98

\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

Log	#	Request	Name	Log	туре	Char	ge Num	Log	Approval	Info
I				L						
đ	Lab		Meth	a Ana	. <b>-</b>					
x	ID		#	b lys	t Analyte	ARL	Re	sult		

96	0928-1	WL-133CONDC	RCRA	522020702 Not Approved
1	6CX56	57012	7RAH ACID	A 5.274E-01 +- 8.6E-03 Normal Acid
2	6CX56	57012	7RAH ACID	A 5.255E-01 +- 8.6E-03 Normal Acid
3	6CX57	87100	7BCS ALUMINUM	
			BORON	A < 1.16162E-03 MOLAR
			IRON	A 6.8E-04 +- 2.3E-04 MOLAR
			ZIRCONIU	M A < 1.00485E-03 MOLAR
			CALCIUM	A 1.17E-03 +- 1.2E-04 MOLAR
4	6CX56	11023	1BCS AL/F RAT	IO A Ratio Not Performed
5	6CX56	57171	7RAH CHLORIDE	A < 2.55951E+01  ug/ml
6	6CX56	57093	7RAH FLUORIDE	C Canceled Entry
7	6CX58	17985	7BCS FLASH PC	IN A NO FLASH @ 60.00 deg C corrected
8	6CX58	17985	7BCS FLASH PC	IN A NO FLASH @ 60.00 deg C corrected
9	6CX56	87970	7BCS GROSS BE	TA C Canceled Entry
10	6CX56	17802	7BCS MERCURY	C Canceled Entry
11	6CX56	97168	7BCS SULFATE	A < 1.84443E+01 uG/mL
12	6CX56	67920	7RLC URANIUM	A < 2.4629E-03 G/L
13	6CX56	67920	7RLC URANIUM	A < 2.4629E-03 G/L
14	6CX56	7972	7 UDS	C Canceled Entry
15	6CX57	22800	2SDN SODIUM	A 3.1077E+02 ug/mL
			POTASSIU	M A 5.6475E+01 ug/mL
16	6CX56	97074	7BCS NITRATE	C Canceled Entry
17	6CX56	13202	3IDG PU239	A $5.7E+01 +- 1.2E+01 d/s/ml$
			PU238	A $2.40E+03 +- 2.6E+02 d/s/ml$
18	6CX56	13993	3MLE CS137	A $1.025E+05 +- 7.1E+03 d/s/ml$
			EU154	A $6.79E+02 +- 6.3E+01 d/s/ml$
			NB94	A $3.07E+02 +- 3.2E+01 d/s/ml$
			RU106	A $2.15E+03 +- 2.1E+02 d/s/ml$
			SB125	A $9.79E+03 +- 4.9E+02 d/s/ml$
19	6CX56	13011	3WDT TRITIUM	A 5.09528E+03 +- 1.2E-01 D/S/ML
20	6CX56	13011	3WDT TRITIUM	A 5.25961E+03 +- 1.3E-01 D/S/ML

Log	#	Request Name	Log	g Type	(	Charge Num Log Approval Info
d x		Meth #	a Ana	a- st Analyte	ARL	. Result
21	6CX56	23381	_	TOTAL SR	A	
22	6CX56	3539	3	IODINE	c	
23	6CX56	57017	7RAH		A	<del>-</del>
24	6CX56	57017	7RAH		Α	<del></del>
25	6CX57	32900	7CBG	ANTIMONY	A	
				ARSENIC	A	
				BARIUM	A A	_
				BERYLLIUM		
				CADMIUM	A	
				CHROMIUM LEAD	A A	
				NICKEL	A	
				SELENIUM	A	
				SILVER	A	
				THALLIUM	A	
26	6CX57	32900	7CBG	ANTIMONY	Α	
20	OCAS /	32300	, CDG	ARSENIC		Not Detected: MDL= 795.0 ug/L
				BARIUM	Α	
			,	BERYLLIUM	Α	
				CADMIUM	Α	_
				CHROMIUM	Α	
				LEAD	Α	
				NICKEL	Α	1.3475E+03 ug/L
				SELENIUM	Α	8.65E+02 ug/L
				SILVER	Α	Not Detected: MDL= 37.5 ug/L
				THALLIUM	Α	3.39E+03 ug/L
27	6CX61	32900	7CBG	ANTIMONY	Α	5.6E+01 ug/L
				ARSENIC	Α	•
				BARIUM	Α	<del>_</del>
				BERYLLIUM	Α	<u> </u>
				CADMIUM	Α	
				CHROMIUM	Α	<del>_</del>
				LEAD		Not Detected: MDL= 39.0 ug/L
				NICKEL	A	
				SELENIUM	A A	
				SILVER THALLIUM		Not Detected: MDL= 129.0 ug/L
28	6CX57	12800	2 CDM	MERCURY	A	
29	6CX57	12800		MERCURY	A	~
30	6CX61	12800		MERCURY		Not Detected: MDL= 2.0 ug/L
31	6CX58	8060	8	TOC	c	
32	6CX58	18060	8BGP		A	
J &	301130	10000		TOC(r)	C	<del>-</del>
33	6CX62	18060	8BGP			Not enough sample to run a dup.
34	6CX59	19260	9HCJ			SEE ATTACHED
35	6CX59	19260	9HCJ			SEE ATTACHED
36	6CX60	19260	9HCJ		Α	
37	6CX58	19270		SVOA		- 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1
38	6CX58	19270		SVOA		

Log	#	Request Name	Log Ty	рe	Charge Num Log Approval Info
I			L		
đ	Lab	Meth	a Ana-		
x	ID	#	b lyst A	nalyte ARI	L Result
39	6CX62	19270	9JXJ SVO	A	. Data Not Approved Yet
40	6CX56	87980	7BCS SPG		~ ~
41	6CX56	87980	7BCS SPG		. 1.02496E+00 @ 25/4
42	6CX56	87980	7BCS SPG		. 1.02496E+00 @ 25/4
43	6CX56	87970		SS BETA A	
44	6CX56	97074	7BCS NIT		_
45	6CX56	57093	7RAH FLU		
46	6CX56	57093	7RAH FLU		
47	6DM47	32900	7CBG ANT		
			ARS	ENIC C	. Canceled Entry
			BAR	IUM C	. Canceled Entry
			BER	YLLIUM C	. Canceled Entry
			CADI	MIUM C	. Canceled Entry
			CHR	OMIUM C	. Canceled Entry
			LEA	C.,	. Canceled Entry
			NIC	KEL C	. Canceled Entry
			SEL	ENIUM C	
			SIL	JER C	. Canceled Entry
			THAI	LLIUM C	. Canceled Entry

\*\*\*\*\*\*\* END \*\*\*\*\*\*\*



Log Search

```
Run by: JEFF LONG
Date of Search: 2003-06-16 14:07:00.293
*************
Search Criteria:
  Start Log....:800101 1
  End Log....:801231 15
  Log Approval.:ALL Logs
  Result Type..:All Entries
  Lab/Group....:ALL Groups
  Name Column..:Lab Sample ID
    Request Name..:*WL-102*
Total # Logs Found...: 1
Total # Results Found: 12
********************
        Request Name Log Type Charge Num Log Approval Info
                     L
 I
 d Lab
                  Meth a Ana-
                  # b lyst Analyte ARL
                                              Result
 x ID
                                      13301-240-412Unapproved by
800616-9 WL-102
               5971 CLD GROSS-B ... +- 1.34437E+06+-41010.6 B/MIN/ML
34 WL-102
35 WL-102
                3993 LEE GAMMA SCAN ... ATTACHED
36 WL-102
               11000 JGJ SODIUM ... 4.32 & 4.34 G/L
37 WL-102
               11000 JGJ POTASSIUM ... 1.04 & 1.02 G/L
38 WL-102
                5171 JMR CHLORIDE ... +- 126.404+-8.97287 UG/ML
39 WL-102
                5092 BLH FLUORIDE ... +- 127.292+-5.89728 UG/ML
                                   ... +- .736395+-2.93917E-02 MOLAR
                5071 NWH NITRATE
40 WL-102
                     HAS SEMI-QUANT ... ATTACHED
41 WL-102
               11000
                     BLH ACID ... +- .473127+-2.07193E-02 NACID
42 WL-102
                5015
                                  ... +- 1.0297+-4.28787E-04
                6981 CLD SP-GR ...
5961 DBB NA/K ...
43 WL-102
66 WL-102
                5961 DBB SEMI-QUANT ...
67 WL-102
```

\*\*\*\*\*\*\* END \*\*\*\*\*\*\*



# EXON NUCLEAR IDAHO COMPANY, Inc.

Internal Correspondence

Date: October 5, 1983

To: W. B. Palmer WBP

From: D. W. Rhodes/Rhod-17-83 Will Rhodes

Subject: Composition of PEW Eyaporator Feed, Condensate and Bottoms

#### Distribution:

- D. R. Alexander
- B. R. Dickey
- G. W. Hogg
- J. E. Johnson
- G. E. Lohse
- L. W. McClure
- W. A. Mickelson
- R. E. Mizia
- E. P. Mondok
- P. I. Nelson
- A. P. Roeh
- R. E. Schindler
- M. D. Staiger
- M. C. Swenson
- F. S. Ward
- D. W. Rhodes-2

During the period August 5 through August 11, 1983, a sampling program was completed to help define the concentrations of chemicals and radionuclides in PEW feed, condensate and bottoms while the NWCF was operating.\* The results of similar sampling programs were reported earlier in (1) Rhod-6-81 (uranium recovery process operating) and (2) Rhod-1-81 (WCF operating).

The feed solution to the evaporator during the test consisted of waste from tanks WG-100(2), WH-101, WC-119, NCD-123, WL-104, and SFE-106. The waste solutions from the PEW-CFD tanks, WG-100 and WH-101, were analyzed as a composite. The condensate samples from WL-107(2), WL-163, and WL-106 were also analyzed as a composite. The results are given in Tables I, II, and III. No interpretation of the results was attempted, since each user has a different need. If you have any questions, please contact me at 6-3080.

<sup>\*</sup>The uranium extraction systems were not operating and the Rover burner operated only one day during the sampling period.

Table I

Chemical Composition of PEW Evaporator
Feed Solutions, Condensate and Bottoms

				٧	essel Num	ber		
Component	<u>Unit</u>	WG-100 WH-101	WC-119	NCD 123	WL-104	SFE-106	WL-613	WL-106 WL-107 WL-163
C1 F S04 Hg A1 B Car Fe Mn Mo Nai Pb K Tin NO3 pH	mg/L	67 16 83 0.2 240 ND 11 ND ND 19 ND ND 620 ND ND 20 23 ND ND 13640	135 29 12 6.8 9.6 3.9 37 0.6 3.4 8.4 0.3 ND 10 0.1 ND 3.8 3 ND 120646 0.77	97 296 42 4E-3 14 ND 230 52 39 160 1500 ND 6110 ND 21 71 1927 12 ND 5580 12.5	83 1424 104 0.7 460 20 160 2.6 13 55 0.6 6.6 185 ND 35 44 79 3.3 ND 4588 1.48	195 0.9 24 2E-3 100 ND 250 ND 16 27 2.4 ND 488 1.4 ND 29 8 ND ND 220 2.34	2435 1777 6336 1201 2700 1800 18000 580 1100 1800 7000 ND 110000 48 470 430 26000 ND 39 556000	43 47 7.7 ND 2 1 0.4 0.7 0.3 ND ND 0.2 ND ND ND 2 2 1 1 1 1 1 1 1 1 1 1 1 1 1
ND = Not [	)etecte	<u>a</u>						

Table II

Sources of Waste Solutions During Boildown Test\*

<u>Date</u>	Feed Source	Gallons	<u>Tank</u>	Condensate Batch No.	Gallons
7-5 7-6 7-7 7-7 7-10 7-11	WG-100 SFE-106 WC-119 WH-101 NCD-123 WG-100	2650 1250 2100 2900 2700 2550	WL-107 WL-163 WL-106 WL-107	227 228 229 230	4000 4000 4000 4000 16000
7-11	WL-104 Tota	<u> </u>	WL-613	Evaporator Botto	ms <u>200</u> 16200

<sup>\*</sup>The difference between the feed volume and the volume of condensate plus evaporator bottoms is 16200-14850=1350 gal. This difference is due to the input of small volumes of water from a variety of sources (VOG loop seals, off-gas drains, pump priming, etc.), that are not measured, as well as instrument and/or reading errors.

Table III

Radiochemical Composition of PEW Evaporator
Feed Solutions, Condensate and Bottoms

				٧	essel Num	ıber 🦠		
Component	<u>Unit</u>	WG-100 WH-101	WC-119	NCD 123	WL-104	SFE-106	<u>WL-613</u>	WL-106 WL-107 WL-163
Total U I-129 Ce-144 Co-60 Cs-137 Eu-152 Eu-154 Eu-155 Ru-106 Sr-90 Sb-125 Mn-54 Pr-144 Y-90 H-3 Nb-95 K-40 Zr-95 ND = Not D	mg/L μCi/mL	6.1E-3 3.2E-4 6.3E-4 2.6E-2 ND ND 0.29 3.3E-2 6.1E-3 ND 6.1E-3 3.3E-2 1.7E-3 ND ND	<0.055 1.7E-4 9.8E-2 5.5E-5 1.5E-2 1.6E-1 ND 1.1E-3 6.0E-4 1.0E-2 0.10 1.8E-3 ND 9.8E-2 0.10 0.26 5.6E-4 ND 1.0E-3	<0.055 3.1E-6 6.5E-5 6.6E-4 ND 5.7E-4 ND ND 2.3E-3 7.7E-6 5.7E-5 ND 6.5E-5 7.7E-6 1.5E-4 2.1E-5 ND ND	3.9 5.5E-6 ND 1.5E-4 7.1E-4 1.9E-2 ND ND 5.9E-4 1.9E-2 3.3E-3 3.9E-2 4.9E-5 ND 3.3E-3 ND ND ND	0.22 2.9E-5 5.8E-4 6.3E-4 ND 3.7E-4 1.8E-2 1.4E-2 3.1E-3 ND ND 1.2E-2 ND ND 5.8E-4 1.2E-2 3.6E-4 ND	217 2.1E-4 1.2 2.7E-2 0.17 2.73 0.16 0.13 4.7E-2 3.9 3.5 0.45 ND 1.2 3.5 2.1E-2 9.7E-2 ND 8.6E-2	<0.055 8.2E-6 5.4E-7 ND 2.8E-7 5.2E-6 ND ND 4.3E-5 1.9E-6 4.1E-6 ND 5.4E-7 1.9E-6 4.0E-2 8.2E-7 ND ND

## **COVER PAGE**

1. SDG Transmittal Date: 5/15/01	
Subcontractor Name: INTEC ANALYTICAL Cl Analytical Laboratories De Bechtel BWXT Idaho, LLC	partment
3. Contract Number: ER-SOW-169	. # . #
4. SDG Type: Volatile Organics by GC/MS	·
5. Reporting Tier: Tier II	Driginal Epidological Epidological WM-1810-010507-05 Entired
6. <u>SDG Number:</u> WM-186-010307	EP1006 000 7 - 02 WM-186-01050 7 - 02 WM-186-01050 7 - 02
7. SAP Number: N/A	Enteres
8. Applicable TOS Modification Numbers: N/A	
1 State	5/15/01
Jeffrey L. Jeter Acting ALD Organic Analyses Supervisor	Date
Shelly J. Sailer	5/15/0 / Date
ALD Quality Assurance Officer	
Jeffrey L. Jeter	_ <i>5/15/0/</i> Date
Volatile Analyses Technical Leader	ORIGINAL

#### TOTAL VOC ANALYSIS DATA SHEET

## Idaho National Engineering and Environmental Laboratory Analytical Laboratories Department

Lab Code:

ALDINTEC

Contract Number:

NΑ

TOS Number:

SDG Number:

WM-186-010307

ALD Report Number: 0103072

Analytical Batch No: 0103072

Field Sample ID:

Method Number:

9260

Matrix:

WATER

Lab Sample ID:

1AM76

Sample wt/voi:

5 mL

WM-186-010307-TB

Lab File ID:

SV102644

Level (low/med):

Low

Date Sampled:

03/07/2001

%Solid:

0

Date Received:

03/08/2001

GC Column:

VOCOL

0.250 (mm)

Date Extracted:

03/21/2001

Soil Extract Volume:

N/A

Date Analyzed:

Soil Aliquot Volume:

N/A

Dilution Factor:

1

ell Erland orative or

		CONCENTRATION	·
CAS#	COMPOUND	(ug/L)	Q
74-87-3	Chloromethane	10	·U
75-01-4	Vinyi Chloride	.10	U
74-83-9	Bromomethane	10	U
75-00-3	Chloroethane	10	U
75-69-4	Trichlorofluoromethane	10	U
75-35-4	1,1-Dichloroethene	10	U·
76-13-1	1,1,2-Trichloro-1,2,2-trifluoroethane	10	U
75-15-0	Carbon Disulfide	10	U
67-64-1	Acetone	20	UZY
75-09-2	Methylene Chloride	20	UZY
156-60-5	trans-1,2-Dichloroethene	10	U
75-34-3	1,1-Dichloroethane	10	U
156-59-2	cis-1,2-Dichloroethene	10	U
78-93-3	2-Butanone	10	U.
67-66-3	Chloroform	10	U
71-55-6	1,1,1-Trichloroethane	10	U
56-23-5	Carbon Tetrachloride	10	UM
71-43-2	Benzene	10	U
107-06-2	1,2-Dichloroethane	20	UMZ
79-01-6	Trichloroethene	10	U

05/14/2001

Prog. Ver.: 1.0

FORM I VOC - RCRA

Page 1 of 2

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019/45

## TOTAL VOC ANALYSIS DATA SHEET

# Idaho National Engineering and Environmental Laboratory Analytical Laboratories Department

Lab Code:

ALDINTEC

Contract Number:

NA

TOS Number:

NA

SDG Number:

WM-186-010307

ALD Report Number: 0103072 Field Sample ID:

WM-186-010307-TB

Analytical Batch No: 0103072

9260

Matrix:

WATER

Method Number:

Sample wt/vol:

Lab Sample ID:

1AM76

5 mL

Lab File ID:

SV102644

Level (low/med):

Low

Date Sampled:

03/07/2001

%Solid:

0

Date Received:

03/08/2001

GC Column: Soil Extract Volume: VOCOL

0.250 (mm)

Date Extracted:

03/21/2001

Soil Aliquot Volume:

N/A

Date Analyzed:

(A)

N/A

Dilution Factor:

CAS#	COMPOUND	CONCENTRATION (ug/L)	Q
78-87-5	1,2-Dichloropropane	10	U
75-27-4	Bromodichloromethane	10	lu
10061-01-5	cis-1,3-Dichloropropene	10	U
108-10-1	4-Methyl-2-Pentanone	20	UZ
108-88-3	Toluene	10	U
10061-02-6	trans-1,3-Dichloropropene	10	U
79-00-5	1,1,2-Trichloroethane	10	U
127-18-4	Tetrachloroethene	10	U
591-78-6	2-Hexanone	20	UMZ
124-48-1	Dibromochloromethane	10	U
108-90-7	Chlorobenzene	10	U
100-41-4	Ethylbenzene	10	U
108-38-3	m-xylene and 106-42-3 p-xylene	20	U
95-47-6	o-Xylene	10	U
00-42-5	Styrene	10	U
5-25-2	Bromoform	10	U
9-34-5	1,1,2,2-Tetrachloroethane	20	UMZ

05/14/2001

Prog. Ver.: 1.0

FORM I VOC - RCRA

Page 2 of 2

**REV 10/98** 

020/45

## TOTAL VOC ANALYSIS DATA SHEET TENTATIVELY IDENTIFIED COMPOUNDS

Idaho National Engineering and Environmental Laboratory Analytical Laboratories Department

Lab Code:

ALDINTEC

Contract Number:

NA

TOS Number:

NA

SDG Number:

WM-186-010307

ALD Report Number: 0103072

Analytical Batch No: 0103072

Field Sample ID:

WM-186-010307-TB

Method Number:

9260

Matrix:

WATER

Lab Sample ID:

1AM76

Sample wt/vol:

Level (low/med):

5 mL

Lab File ID:

SV102644

Low

Date Sampled:

03/07/2001

%Solid:

0

Date Received:

03/08/2001

GC Column: Soil Extract Volume: VOCOL N/A ID:

0.250 (mm) Date Extracted: Date Analyzed:

03/21/2001

Soil Aliquot Volume:

N/A

Dilution Factor:

Number of TICs Found: 0

			Concentration		Retention
	CAS Number	Tentatively Identified VOCs	(ug/L)	Q .	Time (min)
1		Noņe			

FORM I-TIC VOC - RCRA 021/16

## COVER PAGE

1.	Transmittal Date:	04/26/01	
2.	Subcontractor Name:	INTEC Analytical Chemistry Laboratory Analytical Laboratories Department Bechtel BWXT Idaho, LLC	
3.	Contract Number:	Not Applicable	
4.	SDG Type:	Type- 1B	
5.	Reporting Tier:	Tier-2	
6.	Delivery Schedule:	Not Applicable	
7.	SDG Number:	WM-186-010307	2
8.	LTI Number:	01-03072	
9.	TOS Number:	Not Applicable	-
10.	TOS Modification Number:	Not Applicable	
11.	IDP Cost:	Not Applicable	
12.	FSID Numbers:	WM-186-010307	· Metals
13.	Billable OC Samples:	None Original	<del>-</del>
		Original 306: WM-1810-010507 Scaple: BF100160101	

ORIGINAL

Lab Name: INEEL - ACL

Lab Code: ALDCPP Contract:BOP

TOS # : NA

LTI # : 0103072

SDG # :WM-186-010307

Matrix : WATER

Date Received: 03/07/2001

Field Sample ID#:

WM-186-010307

Lab Sample ID#: 1AM75

Sample Preparation Method(s) SEE NARRATIVE

% Solids: 0.0 (~~)

Concentration Units (ug/L or mg/kg wet weight basis): UG/L

			· 1	1	i	
CAS No.	Analyte	Concentration	С	Q.	M	v
7429-90-5	ALUMINUM	6.87E+06			P	
7440-36-0	ANTIMONY	2.73E+02	ש		P	
7440-38-2	ARSENIC	5.45E+02	В		₽	
7440-39-3	BARIUM	3.03E+03			P	
7440-41-7	BERYLLIUM	4.04E+01	В	i	P	
7440-43-9	CADMIUM	1.39E+05	_		P	l
7440-47-3	CHROMIUM	2.16E+05			P	
7440-48-4	COBALT	1.20E+03			₽	
7440-50-8	COPPER	4.08E+04	_		P	
7439-92-1	LEAD	1.02E+05	_	•	P	
7439-96-5	MANGANESE	6.85E+05	_		₽	<b> _</b> _
7439-97-6	MERCURY	2.60E+05	1_1		CV	<u> </u>
7440-02-0	NICKEL	8.99E+04	<b> </b> _		P	<u> </u>
7782-49-2	SELENIUM	3.03E+02	В		P	<u> </u>
7440-22-4	SILVER	2.83E+02	В		P	<u> </u>
7440-28-0	THALLIUM	3.84E+02	ש		P	<u> </u>
7440-62-2	VANADIUM	5.96E+02	В		P	
7440-66-6	ZINC	2.59E+04	1_		P	
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Cas No. Chemical Abstracts Service Registry Number; C = concentration qualifier code; Q = data qualifier Code;

M = method qualifier code; V = validation qualifier Code;

Comments:	
	•

SDG TYPE-1B FORM #1

SOW156

Co153.101.20

Original

WM-186-0103072-1

WM-186-0103072-1

WM-186-0103072-1

Enduct 10/4/2001

Enduct 10/4/2001

## DATA REPORT COVER PAGE

1.	Transmittal Date:	May 31, 2001
2.	Subcontractor Name:	INTEC Analytical Chemistry Laboratory  Analytical Laboratories Department  BECHTEL BWXT LLC, Idaho
3.	Contract Number:	PLN-613
4.	SDG Type:	Type – 3
5.	Reporting Tier:	Tier – 2
6.	Delivery Schedule:	Not Applicable
7.	SDG Number:	WM-186-0103072
8.	LTI Number:	0103072
9.	TOS Number:	Not Applicable
10.	TOS Modification Number:	Not Applicable
11.	IDP Cost:	Internal BBWI charge number credited with all applicable work hours performed.
12.	FSID Numbers:	WM-186-010307
		· · · · · · · · · · · · · · · · · · ·
13.	Billable QC Samples:	None

ORIGINAL

MISCELLANEOUS CLASSICAL ANALYSIS DATA SHEET

Page 1 of 1

Lab Name: INEEL - ACL

Lab Code: ALDCPP

Contract: ER SOW

TOS # : NA

LTI # : 0103072

SDG # :WM-186-010307 Z

Matrix : WATER

Date Received: 03/08/2001

5-30-0

Field Sample ID#:

WM-186-010307

Lab Sample ID#: 1AM75

% Solids: N/A

Analyte	Concentration	UNITS	С	Q	METHOD	v
ACID  FLUORIDE  TIC  TOC  UDS	1.5  2970  119  562	N ACID MG/L MG/L MG/L MG/L	ם ם –	E	ACMM 7012 ACMM 7093 ACMM 8060 USEPA 9060 ACMM 7972	
URANIUM	101	MG/L	-		ACMM 7920	  
			-  -  -  -			
			-  -  -			
			-  -  -			

C =	concentration	qualifier	code;	Q =	data	qualifier	Code;
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Comments:

SDG TYPE-3 FORM #1

SOW156

V = validation qualifier Code;

#### FINAL REPORT for WM-186 SOLIDS

Log Type: \*\* TECHNICAL \*\*

Log Number : 01-05245 Phone Number : 6-7196 Report for : T.G. GARN Mailstop : 5218

Date Approved : Feb 11 2002 Time Approved : 10:26 Date Received: May 24 2001

Time Received: 16:26

GWA charged : 561DTA101 Reviewed by TIFFANY PARK

MSA mR/hr : WARM Signature \_\_\_\_\_

Hazard Index : 1E4 Laboratory QA Review

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS:

	Lab Field	
Analysis	Spl ID Spl ID	Method Analyst Results
Aluminum	1BF85 WM-186 SOLID LIB	42900 RHH 2.23659E+00 wt%
Am241	1BF85 WM-186 SOLID LIB	13201 IDG 4.04E+03 +- 4.0E+02 d/s/g
Antimony	1BF85 WM-186 SOLID LIB	42900 RHH Not Detected: IDL= 0.002292683 w
Arsenic	1BF85 WM-186 SOLID LIB	42900 RHH Not Detected: IDL= 0.001414634 w
Barium	1BF85 WM-186 SOLID LIB	42900 RHH 1.5122E-03 wt%
Beryllium	1BF85 WM-186 SOLID LIB	42900 RHH 4.87805E-05 wt%
Boron	1BF86 WM-186 SOLID NAOH 1BF85 WM-186 SOLID LIB	42900 RHH 2.11091E-02 wt%
CURIUM-242	1BF85 WM-186 SOLID LIB	13960 IDG 3.19E+00+-2.10E+00 d/s/g
Cadmium	1BF85 WM-186 SOLID LIB	42900 RHH 4.06341E-02 Wt%
Calcium	1BF85 WM-186 SOLID LIB	42900 RHH 4.67976E-01 wt%
Cerium	1BF85 WM-186 SOLID LIB	42900 RHH Not Detected: IDL= 0.002780488 w
Cesium	1BF85 WM-186 SOLID LIB	12800 SDN <.00114 wt%
Chloride	1BF87 WM-186 SOLID NACO	28202 NWJ 1.58766E+03 ug/g
Chromium	1BF85 WM-186 SOLID LIB	42900 RHH 7.78537E-02 wt%
Co60	1BF85 WM-186 SOLID LIB	33993 SJH 8.56E+04 +- 7.3E+03 pC/g
Cobalt	1BF85 WM-186 SOLID LIB	42900 RHH Not Detected: IDL= 0.0004878049
Copper	1BF85 WM-186 SOLID LIB	42900 RHH 1.81951E-02 wt%
Cs134	1BF85 WM-186 SOLID LIB	33993 SJH 1.12E+05 +- 1.0E+04 pC/g
Cs137	1BF85 WM-186 SOLID LIB	33993 SJH 9.09E+07 +- 2.6E+06 pC/g
Curium-244	1BF85 WM-186 SOLID LIB	13960 IDG 3.45E+01+-8.93E+00 d/s/g
Eu154	1BF85 WM-186 SOLID LIB	33993 SJH 2.28E+05 +- 3.3E+04 pC/g
Fluoride	1BF87 WM-186 SOLID NACO	28201 NWJ 6.23088E+02 ug/g
Gadolinium	1BF85 WM-186 SOLID LIB	
Iron	1BF85 WM-186 SOLID LIB	
Lead	1BF85 WM-186 SOLID LIB	42900 RHH 3.03415E-02 wt%
Lithium	1BF86 WM-186 SOLID NAOH	42900 RHH 6.0E-04 wt%
Magnesium	1BF85 WM-186 SOLID LIB	42900 RHH 1.71024E-01 wt%
Manganese	1BF85 WM-186 SOLID LIB	42900 RHH 1.87171E-01 wt%
Mercury	1BF85 WM-186 SOLID LIB	12800 SDN < 0.00013 Wt%
Molybdenum	1BF85 WM-186 SOLID LIB	42900 RHH 1.29268E-02 wt%
NB94	1BF85 WM-186 SOLID LIB	33993 SJH 2.77E+04 +- 2.1E+03 pC/g
Nickel	1BF85 WM-186 SOLID LIB	42900 RHH 5.37073E-02 wt%
Niobium	1BF85 WM-186 SOLID LIB	42900 RHH 2.57561E-02 wt%
Nitrate	1BF86 WM-186 SOLID NAOH	28204 NWJ 2.46185E+05 ug/g

#### FINAL Report for 01-05245 continued

	FINAL Report fo	r 01-05245 continued
Analysis	Spl ID Spl ID	Method Analyst Results
Np237	1BF85 WM-186 SOLID LIB	
PHOSPHORUS	1BF85 WM-186 SOLID LIB	42900 RHH 2.4022E+00 wt%
PU238	1BF85 WM-186 SOLID LIB	13202 IDG 1.62E+05 +- 1.5E+04 d/s/g
PU239	1BF85 WM-186 SOLID LIB	13202 IDG $2.71E+04 +- 2.6E+03 d/s/g$
Palladium		42900 RHH 1.74634E-02 wt%
Phosphate	1BF87 WM-186 SOLID NACO	28203 NWJ 5.51802E+04 ug/g
Potassium	1BF85 WM-186 SOLID LIB	12800 SDN 1.63 WT%
Ruthenium	1BF85 WM-186 SOLID LIB	42900 RHH Not Detected: IDL= 0.001512195 w
Sb125	IDIOS WE TOO DODID DID	JJJJ Bon 2.37E.03 . 1.0E.04 pc/g
Selenium	1BF85 WM-186 SOLID LIB	42900 RHH Not Detected: IDL= 0.002341463 w
Silicon	1BF85 WM-186 SOLID LIB	
Silver	1BF85 WM-186 SOLID LIB	
Sodium	1BF85 WM-186 SOLID LIB	
Strontium	1BF85 WM-186 SOLID LIB	
Sulfate	1BF86 WM-186 SOLID NAOH	
Sulfur	1BF85 WM-186 SOLID LIB	
TCLP SAMPLE PREP		17998 BGP Done 6/12/01. Store at RAL stati
Thallium	1BF85 WM-186 SOLID LIB	
Tin	1BF85 WM-186 SOLID LIB	
Titanium	1BF85 WM-186 SOLID LIB	
Total Sr	1BF85 WM-186 SOLID LIB	23381 PAT 1.83E+05 +- 2.7E+04 D/S/G
Tritium	1BF85 WM-186 SOLID LIB	
U234	1BF85 WM-186 SOLID LIB	
U235	1BF85 WM-186 SOLID LIB	
U236	1BF85 WM-186 SOLID LIB	13209 IDG 1.23E+01+-6.46E+00 D/S/G
U238	1BF85 WM-186 SOLID LIB	13209 IDG 5.59E+00+-7.40E+00 D/S/G
Uranium	1BF85 WM-186 SOLID LIB	
Vanadium	1BF85 WM-186 SOLID LIB	
Zinc	1BF85 WM-186 SOLID LIB	
Zirconium	1BF85 WM-186 SOLID LIB	42900 RHH 1.91732E+00 wt%
End of Report	64 results.	



Date of Search: 2003-06-16 14:21:46.965 Run by: JEFF LONG \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* Search Criteria: Start Log....:961008 15 End Log.....:961008 15 Log Approval.:ALL Logs Result Type..:All Entries Lab/Group....:ALL Groups Name Column..:Lab Sample ID Total # Logs Found...: 1 Total # Results Found: 9 \* Log # Request Name Log Type Charge Num Log Approval Info L I d Lab Meth a Ana-# b lyst Analyte ARL Result x ID 961008-15 WL-103 522020702 961012 16:48 CLAYNE B GRIGG PLANT 5 6 7

\*\*\*\*\*\* END \*\*\*\*\*\*

8

9 6DC01

Date of Search: 2003-06-16 14:22:11.754 Run by: JEFF LONG \* Search Criteria: Start Log....:961214 1 End Log.....:961214 1 Log Approval.:ALL Logs Result Type..:All Entries Lab/Group....:ALL Groups Name Column..:Lab Sample ID Total # Logs Found...: 1 Total # Results Found: 12 \* Log Type Charge Num Log Approval Info Request Name Log # I T. d Lab Meth a Ana-# b lyst Analyte ARL x ID Result 522020702 970415 21:54 ALAN W OLAVESON 961214-1 WL-103 PLANT 87802 7KFM MERCURY A.. Not Detected: MDL=0.009 uG/mL 1 6EA76 2 6EA76 17929 7SRT URANIUM PR A.. 2.0E+00 ml 3 6EA76 87100 7RLC ALUMINUM A.. 2.43E-03 +- 4.4E-04 MOLAR 24900 4DDJ URANIUM. A.. 2.2E-02 g/kg 4 6EA76 U234 A.. 7.0E-03 WEIGHT % U235 A.. 7.2E-01 WEIGHT %
U236 A.. 7.0E-03 WEIGHT %
U238 A.. 9.93E+01 WEIGHT %

87970 7RLC GROSS BETA A.. 5.0E+02 +- 1.1E+02 B/Min/ml 5 6EA76 47981 7RLC SPGR A.. 9.98734E-01 +- 2.6E-04 17920 7KFM URANIUM A.. 2.69E-02 +- 1.8E-03 G/L 6EA76 9.98734E-01 +- 2.6E-04 @ 25/4 6 7 6EA76 17920 7KFM URANIUM A.. 2.65E-02 +- 1.8E-03 G/L 8 6EA76

\*\*\*\*\*\*\* END \*\*\*\*\*\*\*

Date of Search: 2003-06-16 14:22:33.887 Run by: JEFF LONG \*\*\*\*\*\*\*\*\*\*\*\*\* Search Criteria: Start Log....:961008 16 End Log....:961008 16 Log Approval.:ALL Logs Result Type..:All Entries Lab/Group....:ALL Groups Name Column..:Lab Sample ID Total # Logs Found...: 1 Total # Results Found: 9 \* Charge Num Log Approval Info Log # Request Name Log Type L Ι d Lab Meth a Anax ID # b lyst Analyte ARL Result 522020702 961223 15:11 JACQUIE S JANIBAG 961008-16 WL-104 PLANT 87092 7RAH FLUORIDE A.. 3.49E+01 +- 1.7E+00 ug/ml 1 6DC30 57171 7RAH CHLORIDE A.. < 1.25947E+01 ug/ml 6DC30 97168 7BCS SULFATE A.. < 9.03652E+00 uG/mL 6DC30 17012 7RAH ACID A.. titrated less than .5 ml
47981 7BCS SPGR A.. 9.99012E-01 +- 2.6E-04 @ 25/4
17920 7RAH URANIUM A.. 2.35E-02 +- 1.3E-03 G/L
87017 7RLC PH A.. 2.75E+00 +- 1.6E-01 pH
17920 7RAH URANIUM C.. Canceled Entry
17920 7RAH URANIUM A.. 2.23E-02 +- 1.6E-03 G/L 6DC30 5 6DC30 6 6DC30 6DC30 8 6DC30 6DC30 9

\*\*\*\*\*\*\* END \*\*\*\*\*\*\*

\*\*\*\*\*\*\* END \*\*\*\*\*\*

			Log Search	
		arch: 2003-06-1		Run by: JEFF LONG
			******	*******
Sea	rch Crit	ceria: og:961214 :	)	
		:961214 :		
	_	coval.:ALL Logs	•	
		Type:All Entr	ies	
		ıp:ALL Grou		
	Name Co	lumn:Lab Samp	le ID	
TOF	al # T.o.	gs Found: 1		
		sults Found: 12		
***	****	*****	*****	*******
_	н			C)
Log	#	Request Name	Log Type L	Charge Num Log Approval Info
	Lab	Met]	n a Ana-	
_	ID	#	b lyst Analyte	ARL Result
961	214-2	WL-104	PLANT	522020702 970415 22:30 ALAN W OLAVESON
1	6EA78		7KFM MERCURY	
2	6EA78	87100	7RLC ALUMINUM	A 3.50E-03 +- 4.5E-04 MOLAR
3	6EA78	87970		A 1.20E+03 +- 1.9E+02 B/Min/ml
4	6EA78	24900	4DDJ URANIUM.	A 2.4E-02 g/kg
			U234 U235	A 0.0E+00 WEIGHT % A 7.0E-01 WEIGHT %
			U235	A 4.0E-03 WEIGHT %
			U238	A 9.93E+01 WEIGHT %
5	6EA78	17929		A 2.0E+00 ml
6	6EA78	47981	7RLC SPGR	A 9.98949E-01 +- 2.6E-04 @ 25/4
7	6EA78	17920	7KFM URANIUM	
8	6EA78	17920	7KFM URANIUM	A 2.43E-02 +- 1.7E-03 G/L

\*\*\*\*\*\*\* END \*\*\*\*\*\*

Run by: JEFF LONG Date of Search: 2003-06-16 14:23:17.773 \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* Search Criteria: Start Log....:970916 9 End Log....:970916 9 Log Approval.:ALL Logs Result Type..:All Entries Lab/Group....:ALL Groups Name Column..:Lab Sample ID Total # Logs Found...: 1 Total # Results Found: 10 \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* Log # Request Name Log Type Charge Num Log Approval Info I L d Lab Meth a Anax ID # b lyst Analyte ARL Result 522020602 970929 17:30 ROBERTA A JORDAN 970916-9 WL-105:105 PLANT 87012 7SRT ACID A.. <.0241 Normal Acid 87017 7SRT PH A.. 2.404E+00 +- 5.6E-02 1 7HR36 7HR36 57171 7RAH CHLORIDE A.. < 4.11211E+00 ug/mL 3 7HR36 87092 7RAH FLUORIDE A.. 5.07E-04 +- 8.1E-05 molar 7HR36 5 17920 7KFM URANIUM A.. < 3.46607E-01 ug/mL 7HR36 17920 7KFM URANIUM A. < 3.46607E-01 ug/mL 87100 7BCS ALUMINUM A. < 6.72322E-04 MOLAR 97168 7BCS SULFATE A. < 4.06935E+00 ug/ml 87970 7KFM GROSS BETA A. 2.05E+03 +- 1.9E+02 B/Min/ml 87100 7BCS ALUMINUM A. < 4.74571E-04 MOLAR 6 7HR36 7HR36 8 7HR36 9 7HR36 10 7HR36



```
Log # Request Name Log Type Charge Num Log Approval Info
    I
d Lab
                                  L
                                                         Meth a Ana-
     x ID
                                                           # b lyst Analyte ARL
                                                                                                                                                  Result
930513-29 WM-100 13120-200-001930601 04:14 SPLMGT SPLMGT 75 CANCELLED 1 1 LOG C.. -- No sample received --
                                              7961 7RLC RADIO CHEM C.. Prep Completed 082193 07:33
53993 3DAO GAMMA SCAN C.. ** See Index 478 thru 481. **
23381 3BJS TOTAL SR A.. 5.95137+-.289466 UCI/ML
2110 2JSL SODIUM A.. =0.023 Molarity
2190 2JSL POTASSIUM A.. =0.0018 Molarity
77168 7BKH SULFATE A.. 6.05235E-03+-2.86055E-04 Molar
77074 7SRT NITRATE A.. 4.22397+-.646989 Molar
57171 7BET CHLORIDE C.. < 155.44 UG/ML
7972 7WDT UDS A.. * 3.68 MG/ML
47981 7WDT SP-GR A.. 1.21961+-7.80476E-04 @ 25/4
67015 7BET ACID A.. < 8.97114E-02 NACID
67093 7BGP FLUORIDE A.. 2.20899E-02+-1.89826E-03 MOLAR
7961 7RLC SPEC CHEM C.. Prep Completed 082193 07:33
67920 7BET URANIUM A.. 2.40951E-03+-6.96431E-04 G/L
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM C.. > .190546 MOLAR
7103 7 CADMIUM C.. SEE472
7105 7SDN IRON A.. 1.81135E-02+-5.63841E-04 MOLAR
7110 7SDN ZIRCONIUM A.. 2.67232E-03+-5.75093E-04 MOLAR
7111 7SDN CALCIUM A.. 2.67232E-03+-5.75093E-04 MOLAR
57171 7BET CHLORIDE A.. 25.2599+-4.24004 UG/ML
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM A.. 2.67232E-03+-5.75093E-04 MOLAR
7110 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN METALS-ICP C.. See Individual Elements Below
930820-5 WM-100AT
30 1
30 1
31 1
32 1
33 1
34 1
35 1
36 1
37 1
38 1
39 1
40 1
41 1
42 1
43 1
44 1
45 1
46 1
47 1
48 1
49 1
50 1
330.2
                                                     7100 7SDN METALS-ICP C.. See Individual Elements Below
465 1
                                                     7101 7SDN ALUMINUM A.. 1.29549+-.361913 MOLAR
466 1
                                               7101 7SDN ALUMINUM A.. 1.29549+-.361913 MOLA

7100 7SDN METALS-ICP C.. SEE 472

7103 7SDN CADMIUM C.. < 9.50622E-04 MOLAR

53993 3DAO CO-60 A.. 2060+-228 D/S/ML

53993 3DAO CS-134 A.. 19400+-894 D/S/ML

53993 3DAO CS-137 A.. 514000+-17500 D/S/ML

53993 3DAO EU-154 A.. 3230+-383 D/S/ML
471 1
472 1
478 1
479 1
480 1
481 1
                                               7900 7WDT EPA-TOX C.. See Individual Elements Below 7903 7WDT CADMIUM C.. > 130.628 mg/L
485 1
486 1
                                               7903 7WD1 CADMIDM C... 2150.020 Mg/E

57171 7RLC CHLORIDE A.. 30.9039+-7.96776 UG/ML

2110 2 SODIUM C.. NOT REQUESTED

2190 2 POTASSIUM C.. NOT REQUESTED

7961 7WDT NA/K SPEC C.. Prep Completed 090393 18:57
493 1R
494 PLASTIC1R
495 PLASTIC1R
496 PLASTIC1R
                                                    7900 7WDT EPA-TOX C.. See Individual Elements Below 7903 7WDT CADMIUM A.. 100.222+-7.66378 mg/L
497 1
498 1
930820-13 WM-100
                                                                                                                         13120-200-001940217 08:29 SPLMGT SPLMGT
                              47981 7MLE SP-GR A.. 1.26416+-7.80474E-04 @ 25/4
47981 7MLE SP-GR A.. 1.24535+-7.80487E-04 @ 25/4
47981 7MLE SP-GR A.. 1.26386+-7.80474E-04 @ 25/4
375 1
376 2
377 3
```

Log	#	Request Name	Lo	g Type		Charge Num Log Approval Info
Ĭ			L	5 -1F-		onarge nam bog hpprovar into
đ	Lab	Meth	a An	a-		
×	ID	#	b ly	st Analyte	ARL	Result
378		67015		ACID	Α	.120714+-3.74996E-02 NACID
379		67015		ACID	Α	
380		67920		URANIUM	Α	
381		7972	7RLC		Α	* 2.18 MG/ML
382		57171	7BET	CHLORIDE	Α	
383		77074	7SRT	NITRATE	Α	6.40565+636018 Molar
384		67093	7MLE	FLUORIDE	Α	2.22026E-02+-1.89713E-03 MOLAR
385		77168	7вкн	SULFATE	Α	1.01392E-03+-3.09891E-04 Molar
386		7961	7MLE	RADIOCHEM	С	<u> </u>
	PLASTIC 3			SPECTCHEM	С	Prep Completed 082393 14:45
	PLASTIC 2	7961		SPECTCHEM	С	Prep Completed 082393 14:46
389		83993	3DAO	GAMMA SCAN	С	** See Index 483 thru 484. **
390		23381	3BJS	TOTAL SR	Α	13.1827+434105 UCI/ML
391		3205	3IDG	TRANS-UA	Α	6002.79+-344.485 D/S/ML
392		23011	3JSJ	TRITIUM	Α	.297639+-4.64174E-02 D/SEC/ML
	PLASTIC 1			EPA-TOX	С.,	See Individual Elements Below
	PLASTIC 3		7CBG	ARSENIC		< 3.54948  mg/L
395	PLASTIC 3	L 7902		BARIUM	Α	< 4.57491 mg/L
	PLASTIC 3		7CBG	CADMIUM	Α	62.9984+-3.09222 mg/L
397	PLASTIC 3	1 7904	7CBG	CHROMIUM	Α	32.517+-2.81098 mg/L
398	PLASTIC 1	L 7905		LEAD	Α	40.7849+-3.38658 mg/L
399	PLASTIC 1	L 7906	7CBG	SELENIUM	С	< 3.12083 mg/L
400	PLASTIC 1	L 7907	7CBG	SILVER	C	< 4.04898 mg/L
401	PLASTIC 1	L 7908	7CBG	NICKEL	Α	12.5588+-1.0438 mg/L
402	PLASTIC 2	7900	7CBG	EPA-TOX	С	See Individual Elements Below
403	PLASTIC 2	7901	7CBG	ARSENIC	Α	< 3.54948 mg/L
404	PLASTIC 2	7902	7CBG	BARIUM	Α	< 4.57491 mg/L
405	PLASTIC 2	7903	7CBG	CADMIUM	Α	61.2782+-3.04655 mg/L
406	PLASTIC 2	7904	7CBG	CHROMIUM	Α	33.6403+-2.86527 mg/L
407	PLASTIC 2	7905	7CBG	LEAD	Α	44.3699+-3.53409 mg/L
	PLASTIC 2		7CBG	SELENIUM	С	< 3.12083  mg/L
	PLASTIC 2		7CBG	SILVER	С	< 4.04898 mg/L
	PLASTIC 2		7CBG	NICKEL	Α	13.9321+-1.11811 mg/L
	PLASTIC 1			MERCURY	Α	=1.7E+02 mg/1
	PLASTIC 2		2LBZ	MERCURY	Α	=1.6E+02 mg/1
	PLASTIC 1		2	ARSENIC.		
	PLASTIC 2		2			SEE RAL DATA
	PLASTIC 1		2			SEE RAL DATA
	PLASTIC 2		2	SELENIUM.		SEE RAL DATA
	PLASTIC 1			SODIUM		=0.17 Molarity
	PLASTIC 1	- ·				=0.021 Molarity
419		67016	7KFM			< .5 pH
	RAMP WM10					NO Flash @ 60.0 deg C corrected
	PLASTIC 1					Prep Completed 082393 14:46
	VOA VIAL	•		GCMS-2SRC		
	VOA VIAL			GCMS-2SRC		
	PLASTIC 1					See Individual Elements Below
	PLASTIC 1					> .190546 MOLAR
426	PLASTIC 1	. 7102	7SDN	BORON	Α	< 3.75619E-03 MOLAR

```
Log #
                         Request Name
                                                             Log Type Charge Num Log Approval Info
   I
    d Lab
                                               Meth a Ana-
    x ID
                                                #
                                                           b lyst Analyte ARL
                                                                                                                          Result
                                              427 PLASTIC 1
                                        7105 7SDN IRON A.. .02076+-6.04381E-04 MOLAR
7110 7SDN ZIRCONIUM A.. < 1.56506E-03 MOLAR
7111 7SDN CALCIUM A.. 1.16336E-02+-9.42946E-04 MOLAR
47981 7MLE SP-GR A.. 1.2311+-7.80485E-04 @ 25/4
67016 7KFM PH A.. < .5 pH
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN ARSENIC C.. < 5.59712 mg/L
7906 7SDN SELENIUM C.. < 2.53942 mg/L
7907 7SDN SILVER A.. < 2.45596 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN ARSENIC C.. < 5.59712 mg/L
7907 7SDN SILVER A.. < 2.45596 mg/L
7907 7SDN SELENIUM C.. < 5.59712 mg/L
7907 7SDN SELENIUM C.. < 5.59712 mg/L
7907 7SDN SELENIUM C.. < 2.53942 mg/L
7907 7SDN SELENIUM C.. < 2.53942 mg/L
7907 7SDN SILVER A.. < 2.45596 mg/L
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM A.. 1.08878+-.362405 MOLAR
7100 7SDN METALS-ICP C.. See Individual Elements Below
 428 PLASTIC 1
                                                                                           A.. .02076+-6.04381E-04 MOLAR
 429 PLASTIC 1
 430 PLASTIC 1
 431 4
432 6
452 5
453 5
454 5
455 5
456 6
457 6
458 6
459 6
463 PLASTIC 1
464 PLASTIC 1
                                        7101 75DN ALUMINUM A.. 1.000/07-.502405 Molan

7100 75DN METALS-ICP C.. See Individual Elements Below

7103 75DN CADMIUM A.. < 9.50622E-04 MOLAR

83993 3DAO CS-134 A.. 20600+-1120 D/S/ML

83993 3DAO CS-137 A.. 838000+-31900 D/S/ML
467 PLASTIC 1
468 PLASTIC 1
483 5
484 5
                                        7100 7WDT METALS-ICP C.. See Individual Elements Below 7111 7WDT CALCIUM A.. 1.18246E-02+-9.5268E-04 MOLAR
487 PLASTIC1R
488 PLASTIC1R
                                        77074 7MLE NITRATE
489 3R
                                                                                             A.. 5.78416+-.618707 Molar
                                        7985 7CWL FLASHPOINT C.. Flashed @ 52.0 deg C corrected 7985 7CWL FLASHPOINT C.. Flashed @ 52.0 deg C corrected 7985 7CWL FLASHPOINT A.. NO Flash @ 60.0 deg C corrected 7961 7KFM SP CHEM C.. Prep Completed 090893 23:20
490 BEG KNOWN
491 END KNOWN
492 60C WM100
499 PLASTIC 1
                                     7961 7KFM SP CHEM C.. Prep Completed 090893 23:20
7961 7KFM SP CHEM C.. Prep Completed 090893 23:21
7961 7KFM SP CHEM C.. Prep Completed 090893 23:21
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN ARSENIC A.. < 3.89309 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN ARSENIC A.. < 3.89309 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7906 7SDN SELENIUM A.. < .991639 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7906 7SDN SELENIUM A.. < .991639 mg/L
7900 7SDN SELENIUM A.. < .991639 mg/L
500 PLASTIC 1
501 PLASTIC 2
504 PLASTIC 1R
505 PLASTIC 1R
506 PLASTIC 2R
507 PLASTIC 2R
508 PLASTIC 1R
509 PLASTIC 1R
510 PLASTIC 2R
511 PLASTIC 2R
930820-16 WM-100
                                                                                                      13120-200-001940210 23:33 SPLMGT SPLMGT
331 1
                                          47981 7RLC SP-GR A.. .999071+-7.79144E-04 @ 25/4
332 3
                                           47981 7RLC SP-GR
                                                                                          A.. .999458+-7.79146E-04 @ 25/4
                                        67015 7BGP ACID A.. < 8.83383E-02 NACID 67015 7BGP ACID A.. < 8.97114E-02 NACID
333 2
334 3
335 2
                                        87920 7RLC URANIUM A.. < 4.22281E-04 G/L
336 4
                                          7972 7RLC UDS A.. NO VISIBLE SOLIDS
                                      7772 7REC ODS A.. NO VISIBLE SOLIDS
57171 7BET CHLORIDE A.. < 15.0301 UG/ML
77074 7SRT NITRATE A.. < 1.61E-05 Molar
67093 7BGP FLUORIDE C.. < 1.35096E-03 MOLAR
77168 7MLE SULFATE A < 1.04102E-06 Molar
337 2
338 3
339 2
340 3
                                         77168 7MLE SULFATE A.. < 1.04102E-06 Molar
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Log #
                     Request Name Log Type Charge Num Log Approval Info
     I
d Lab
                                                                    Meth a Ana-
                                                 # b lyst Analyte ARL Result

7961 7RLC RADIO CHEM C.. Prep Completed 082193 04:48
7961 7RLC RADIO CHEM C.. Prep Completed 082193 04:48
7961 7RLC SPEC CHEM C.. Prep Completed 082193 04:48
7961 7RLC SPEC CHEM C.. Prep Completed 082193 04:49
53993 3DAO GAMMA SCAN C.. ** See Index 482. **
23381 3BJS TOTAL SR A.. 3.12763E-05+6-78419E-06 UCI/M
3205 3IDG TRANS-UA A.. < .137597 D/S/ML
23011 3JSJ TRITIUM A.. < .157442 D/SEC/ML
7900 7CEG EPA-TOX C.. See Individual Elements Below
7901 7CEG ARSENIC A.. < .67931 mg/L
7902 7CEG BARIUM A.. < .875559 mg/L
7903 7CEG CADMIUM A.. < .830681 mg/L
7904 7CEG CADMIUM A.. < .830681 mg/L
7905 7CEG LEAD A.. < .573449 mg/L
7907 7CEG SILVER A.. < .774906 mg/L
7908 7CEG NICKEL A.. < .597274 mg/L
7907 7CEG SILVER A.. < .774906 mg/L
2330 2 ARSENIC C.. SEE METHOD 7901
2340 2 SELENIUM C.. SEE METHOD 7906
2110 2JSL SODIUM A.. =7.0E-06 Molarity
2190 2JSL POTASSIUM A.. =1.8E-06 Molarity
67016 7KFM PH A.. 9.56302+-.035681 pH
7985 7SDN FLASHPOINT C.. Flashed @ 41.0 deg C corrected
7961 7RLC SPEC CHEM C.. Prep Completed 082193 04:49
9305 9HCJ GCMS-2SRC C.. NOT REQUIRED
7100 7SDN METALS-ICP C.. SEE Individual Elements Below
7101 7SDN ALUMINUM A.. < 6.56896E-03 MOLAR
7102 7SDN BORON A.. 3.96261E-03+-1.52717E-03 MOLAR
7103 7 CADMIUM C.. SEE 469
7105 7SDN IRON A.. < 2.211E-04 MOLAR
7111 7SDN CALCIUM A.. < 2.54008E-03 MOLAR
7110 7SDN METALS-ICP C.. SEE Individual Elements Below
7101 7SDN METALS-ICP C.. SEE Individual Elements Below
7103 7SDN CADMIUM A.. < 2.54008E-03 MOLAR
7110 7SDN METALS-ICP C.. SEE Individual Elements Below
7101 7SDN METALS-ICP C.. SEE Individual Elements Below
7103 7SDN CADMIUM A.. < 2.54008E-03 MOLAR
7110 7SDN METALS-ICP C.. SEE Individual Elements Below
7103 7SDN CADMIUM A.. < 9.50622E-04 MOLAR
7100 7SDN METALS-ICP C.. SEE Individual Elements Below
7103 7SDN CADMIUM A.. < 9.50622E-04 MOLAR
7100 7SDN METALS-ICP C.. SEE Individual Elements Below
7103 7SDN CADMIUM A.. < 9.50622E-04 MOLAR
7100 7SDN METALS-ICP C.. SEE Individual Elements Below
7103 7SDN CADMIUM A.. < 9.50622E-04 MOLAR
71
       x ID
                                                                                        b lyst Analyte ARL
                                                                                                                                                                                     Result
  341 5
  342 PLASTIC 1
  343 PLASTIC 2
  344 5
 345 5
 346 5
 347 5
 348 PLASTIC 1
 349 PLASTIC 1
 350 PLASTIC 1
 351 PLASTIC 1
 352 PLASTIC 1
 353 PLASTIC 1
 354 PLASTIC 1
 355 PLASTIC 1
 356 PLASTIC 1
 357 PLASTIC 2
 358 PLASTIC 2
359 PLASTIC 1
 360 PLASTIC 1
 361 PLASTIC 1
 362 5
362 5
363 KNOWN
 364 PLASTIC 1
 365 VOA VIAL 1
 366 VOA VIAL 2
 367 PLASTIC 1
 368 PLASTIC 1
 369 PLASTIC 1
 370 PLASTIC 1
 371 PLASTIC 1
 372 PLASTIC 1
 373 PLASTIC 1
 374 2
 434 5
                                                         67016 7KFM PH A.. 8.50498+-3.54009E-02 pH
7100 7SDN METALS-ICP C.. See Individual Elements Below
7103 7SDN CADMIUM A.. < 9.50622E-04 MOLAR
53993 3DAO CS-137 A.. 58.4+-3.39 D/S/ML
 435 6
469 PLASTIC 1
 470 PLASTIC 1
482 5
 502 6
                                                                7985 7SDN FLASHPOINT A.. NO Flash @ 60.0 deg C corrected
 930821-6
                                      WM-100
                                                                                                                                                        13120-200-001940302 15:36 SPLMGT SPLMGT
24 1
25 2
26 3
27 2
28 3
29 2
                                                                 47981 7MLE SP-GR A.. 1.22632+-7.80482E-04 @ 25/4
47981 7MLE SP-GR A.. 1.2212+-7.80478E-04 @ 25/4
                                                              47981 7MLE SP-GR A.. 1.22013+-7.80477E-04 @ 25/4
67015 7BET ACID A.. < 8.97114E-02 NACID
67015 7BET ACID A.. < 8.97114E-02 NACID
67920 7BET URANIUM A.. 3.66657E-03+-8.39782E-04 G/L
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Log I		Request Name	Log Type L		Charge Num Log Approval Info
đ	Lab	Meth	a Ana-		
x		#	b lyst Analyte	ART	Result
					·
30	4	7972	7RLC UDS	Α	* 5.07 MG/ML
31	2	57171	7BET CHLORIDE	С.,	< 120.968 UG/ML
32	3	77074	7SRT NITRATE	Α	
33	2	67093	7MLE FLUORIDE	Α	
34	3	77168	7BKH SULFATE	Α	
35	5	7961	7WDT RADIOCHEM	С	
36	PLASTIC 1	7961	7MLE SPECTCHEM	С	<del>_</del>
37	PLASTIC 2	7961	7MLE SPECTCHEM	С	
38	5	73993			** See Index 279 thru 281. **
39	5	23381	3BJS TOTAL SR	Α	
	5	3205	3IDG TRANS-UA	Α	
41	5	23011	3JSJ TRITIUM		< 157.599 D/SEC/ML
	PLASTIC 1	7900	7CBG EPA-TOX		See Individual Elements Below
43	PLASTIC 1	7901	7CBG ARSENIC		< 3.54948 mg/L
	PLASTIC 1	7902	7CBG BARIUM		< 4.57491 mg/L
	PLASTIC 1	7903	7CBG CADMIUM		22.7578+-1.74276 mg/L
	PLASTIC 1	7904	7CBG CHROMIUM	Α	<del>-</del>
	PLASTIC 1	7905	7CBG LEAD	Α	
	PLASTIC 1	7906	7CBG SELENIUM		< 3.12083 mg/L
	PLASTIC 1	7907	7CBG SILVER		< 4.04898 mg/L
	PLASTIC 1	7908	7CBG NICKEL		4.84096+433302 mg/L
	PLASTIC 2	7900	7CBG EPA-TOX		See Individual Elements Below
	PLASTIC 2	7901	7CBG ARSENIC		< 3.54948 mg/L
	PLASTIC 2	7902	7CBG BARIUM		< 4.57491 mg/L
	PLASTIC 2	7903	7CBG CADMIUM	Α	
	PLASTIC 2	7904	7CBG CHROMIUM		8.02166+-1.09908 mg/L
	PLASTIC 2		7CBG LEAD	Α	
	PLASTIC 2	7906	7CBG SELENIUM		< 3.12083 mg/L
	PLASTIC 2	7907	7CBG SILVER		< 4.04898 mg/L
	PLASTIC 2	7908	7CBG NICKEL		< 4.21435 mg/L
	PLASTIC 1	2809	2LBZ MERCURY	Α	
	PLASTIC 2		2LBZ MERCURY	A	
	PLASTIC 1	2330	2 ARSENIC.	c	
	PLASTIC 2		2 ARSENIC.	c	
	PLASTIC 1		2 SELENIUM.		
	PLASTIC 2				SEE RAL DATA
	PLASTIC 1				=0.049 Molarity
	PLASTIC 1				=0.0046 Molarity
	5				1.67766+-3.60744E-02 pH
	RAMP WM10				NO Flash @ 60.0 deg C corrected
	PLASTIC 1				Prep Completed 082393 14:50
	VOA VIAL				SEE ATTACHED
	VOA VIAL		9HCJ GCMS-2SRC		
	PLASTIC 1				See Individual Elements Below
	PLASTIC 1				> .190546 MOLAR
	PLASTIC 1				< 3.75619E-03 MOLAR
	PLASTIC 1		7 CADMIUM		
	PLASTIC 1				2.02119E-02+-5.96183E-04 MOLAR
	PLASTIC 1		7SDN IRON 7SDN ZIRCONIUM	A	

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Log #
           Request Name
                             Log Type Charge Num Log Approval Info
 I
  d Lab
                      Meth a Ana-
  x ID
                           b lyst Analyte ARL
                                                          Result
79 PLASTIC 1
                           7SDN CALCIUM A.. 3.42787E-03+-6.14036E-04 MOLAR
                     7111
80 4
                    47981
                           7MLE SP-GR
                                           A.. 1.177+-7.80374E-04 @ 25/4
138 3
                    57171 7BET CHLORIDE
                                          A.. 53.9264+-8.5855 UG/ML
244 6
                    67016 7KFM PH
                                           A.. 1.69967+-3.60655E-02 pH
249 PLASTIC 2
                     7961 7MLE SPECTCHEM C.. Prep Completed 082393 14:51
251 60C WM100
                     7985 7CWL FLASHPOINT A.. NO Flash @ 60.0 deg C corrected
252 END KNOWN
                     7985 7CWL FLASHPOINT A.. Flashed @ 60.0 deg C corrected
256 5
                     7900 7SDN EPA-TOX
                                         C.. See Individual Elements Below
257 5
                     7901 7SDN ARSENIC
                                           C.. < 5.59712 \text{ mg/L}
258 5
                     7906 7SDN SELENIUM C.. < 2.53942 mg/L
259 5
                     7907 7SDN SILVER A.. < 2.45596 mg/L
260 5
                     7908 7SDN NICKEL
                                          A.. 5.05519+-.488375 mg/L
261 6
                     7900 7SDN EPA-TOX C.. See Individual Elements Below
262 6
                     7901 7SDN ARSENIC C.. < 5.59712 mg/L
263 6
                     7906 7SDN SELENIUM C.. < 2.53942 mg/L
264 6
                     7907 7SDN SILVER
                                           A... < 2.45596 \text{ mg/L}
265 6
                     7908 7SDN NICKEL
                                           A.. 6.17468+-.591991 mg/L
                     7100 7SDN METALS-ICP C.. See Individual Elements Below
271 PLASTIC 1
272 PLASTIC 1
                     7101 7SDN ALUMINUM A.. 1.04358+-.362522 MOLAR
273 PLASTIC 1
                    7100 7SDN METALS-ICP C.. See Individual Elements Below
274 PLASTIC 1
                    7103 7SDN CADMIUM A.. < 9.50622E-04 MOLAR
                                           A.. 4750+-1310 D/S/ML
279 5
                    73993 3DAO SB-125
280 5
                    73993 3DAO CS-134
                                           A.. 13300+-793 D/S/ML
281 5
                   73993 3DAO CS-137
                                           A.. 415000+-16100 D/S/ML
                   7961 7KFM SP CHEM C.. Prep Completed 090993 20:49
283 PLASTIC 1
                     7961 7KFM SP CHEM C.. Prep Completed 090993 20:49
284 PLASTIC 2
                     7900 7SDN EPA-TOX C.. See Individual Elements Below
293 PLASTIC 1R
                     7901 7SDN ARSENIC A.. < 3.89309 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
294 PLASTIC 1R
295 PLASTIC 2R
                     7901 7SDN ARSENIC A.. < 3.89309 mg/L
296 PLASTIC 2R
297 PLASTIC 1R
                     7900 7SDN EPA-TOX
                                           C.. See Individual Elements Below
298 PLASTIC 1R
                     7906 7SDN SELENIUM A.. < .991639 mg/L
299 PLASTIC 2R
                     7900
                           7SDN EPA-TOX
                                           C.. See Individual Elements Below
300 PLASTIC 2R
                     7906 7SDN SELENIUM A.. < .991639 mg/L
930821-7
           WM-100
                                               13120-200-001940210 23:16 SPLMGT SPLMGT
                                           A.. 1.20464+-7.80453E-04 @ 25/4
A.. 1.20465+-7.80453E-04 @ 25/4
A.. 1.20457+-7.80453E-04 @ 25/4
81 1
                    47981
                           7MLE SP-GR
82 2
                    47981
                           7MLE SP-GR
83 3
                    47981
                           7MLE SP-GR
                                           A.. < 8.97114E-02 NACID
A.. < 8.97114E-02 NACID
A.. < 1.16045E-03 G/L
84 2
                    67015
                           7BET ACID
85 3
                    67015
                           7BET ACID
86 2
                    67920
                           7BET URANIUM
87 4
                                           A.. NO VISIBLE SOLIDS
                     7972
                           7AWO UDS
88 2
                    57171
                           7BET CHLORIDE
                                           A.. < 15.0301 UG/ML
89 3
                                           A.. < 1.61E-05 Molar
                    77074
                           7SRT NITRATE
                                           A.. < 1.33105E-03 MOLAR
90 2
                    67093
                           7MLE FLUORIDE
91 3
                                           A.. < 1.45209E-06 Molar
                    77168 7MLE SULFATE
                   7961 7WDT RADIOCHEM C.. Prep Completed 082393 11:24 7961 7WDT SPEC-CHEM C.. Prep Completed 082393 11:25
92 5
93 PLASTIC 1
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Log	#	Request	Name	Log	g Type		Charge Num Log Approval Info
I				L			
đ	Lab		Meth	a Ana	a-		
x	ID		#	b lys	st Analyte	ARL	Result
	1.1010						
94	PLASTIC	2	7961	7	SPEC PREP	С	NOT REQUESTED
95	5	7	3993	3DAO	GAMMA SCAN	С.,	** See Index 282. **
96	5	2	3381	3BJS	TOTAL SR	Α	<< 5.01646E-06+-3.56361E-06 UCI/M
97	5		3205	3IDG	TRANS-UA	Α	< .137597 D/S/ML
98	5	2	3011	3JSJ	TRITIUM	Α	< 5.6679 D/SEC/ML
99	PLASTIC	1	7900	7CBG	EPA-TOX	С	See Individual Elements Below
100	PLASTIC	1	7901	7CBG	ARSENIC	Α	< .67931 mg/L
101	PLASTIC	1	7902	7CBG	BARIUM	Α	< .875559 mg/L
102	PLASTIC	1	7903	7CBG	CADMIUM	Α	< .863441 mg/L
103	PLASTIC	1	7904	7CBG	CHROMIUM	Α	< .830681 mg/L
104	PLASTIC	1	7905	7CBG	LEAD	Α	< .573449 mg/L
105	PLASTIC	1	7906	7CBG	SELENIUM	Α	< .597274 mg/L
106	PLASTIC	1	7907	7CBG	SILVER	Α	< .774906 mg/L
107	PLASTIC	1	7908	7CBG	NICKEL	Α	< .806555 mg/L
108	PLASTIC	2	7900	7	EPA-TOX	С	NOT REQUESTED
109	PLASTIC	2	7901	7	ARSENIC	С	NOT REQUESTED
110	PLASTIC	2	7902	7	BARIUM	С	NOT REQUESTED
111	PLASTIC	2	7903	7	CADMIUM	С	NOT REQUESTED
112	PLASTIC	2	7904	7	CHROMIUM	С	NOT REQUESTED
113	PLASTIC	2	7905	7	LEAD	С	NOT REQUESTED
114	PLASTIC	2	7906	7	SELENIUM	С	NOT REQUESTED
115	PLASTIC	2	7907	7	SILVER	С	NOT REQUESTED
116	PLASTIC	2	7908	7	NICKEL	С	NOT REQUESTED
117	PLASTIC	1	2809	2LBZ	MERCURY	Α	=Not Detected:DL= 0.0045 mg/l
118	PLASTIC	2	2809	2	MERCURY	С	NOT REQUESTED
119	PLASTIC	1	2330	2	ARSENIC.	С	SEE METHOD 7901
120	PLASTIC	2	2330	2	ARSENIC.	С	NOT REQUESTED
121	PLASTIC	1	2340	2	SELENIUM.	С	SEE METHOD 7906
122	PLASTIC	2	2340	2	SELENIUM.	С	NOT REQUESTED
123	PLASTIC	1	2110	2JSL	SODIUM	Α	=7.3E-05 Molarity
124	PLASTIC	1	2190	2JSL	POTASSIUM	Α	=4.5E-06 Molarity
125	5	6	7016	7KFM	PH	Α	
126	6		7985	7SDN	FLASHPOINT	Α	NO Flash @ 60.0 deg C corrected
127	PLASTIC	1	7961	7WDT	SPEC-CHEM	С.,	Prep Completed 082393 11:26
128	VOA VIAL	. 1	9305				SEE LOG 082016 INDEX 365
129	VOA VIAL		9305	9HCJ	GCMS-2SRC	С	SEE LOG 082016 INDEX 365
130	PLASTIC	1	7100	7SDN	METALS-ICP	С	See Individual Elements Below
	PLASTIC		7101	7SDN	ALUMINUM	Α	< 6.56896E-03 MOLAR
132	PLASTIC	1	7102	7SDN	BORON	Α	< 3.75619E-03 MOLAR
	PLASTIC		7103	7	CADMIUM	С	NOT REQUESTED
	PLASTIC		7105	7SDN			< 2.211E-04 MOLAR
135	PLASTIC		7110				< 1.56506E-03 MOLAR
	PLASTIC		7111				< 2.54008E-03 MOLAR
137	4		7981				1.20458+-7.80453E-04 @ 25/4
250			7961				Prep Completed 082393 11:26
282			3993				14.4+-1.85 D/S/ML
292	KNOWN		7985				Flashed @ 41.0 deg C corrected

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Log # Request Name Log Type Charge Num Log Approval Info
   I
    d Lab
                          Meth a Ana-
   x ID
                           # b lyst Analyte ARL
                                                                    Result
 930822-3
                WM-100
                        13120-200-001940302 15:22 SPI
47981 7MLE SP-GR C. 1.44039+-7.80554E-04 @ 25/4
47981 7MLE SP-GR C. 1.44165+-7.80566E-04 @ 25/4
47981 7MLE SP-GR C. 1.43988+-7.8055E-04 @ 25/4
67015 7MLE ACID A. .116755+-.037468 NACID
67015 7MLE ACID A. .107615+-3.73951E-02 NACID
                                                         13120-200-001940302 15:22 SPLMGT SPLMGT
 930822-3
25 1
26 2
27 3
28 2
29 3
30 2
31 4
32 2
33 3
34 2
                        67920 7BET URANIUM A.. 9.49411E-03+-1.31493E-03 G/L
7972 7RLC UDS C.. * 1.25 MG/ML
                        57171 7MLE CHLORIDE A.. 198.82+-42.284 UG/ML
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Log # Request Name
                                                                                               Log Type Charge Num Log Approval Info
      I
        d Lab
                                                                       Meth a Ana-
       x ID
                                                                           #
                                                                                            b lyst Analyte ARL
                                                                                                                                                                                                Result
73 VOA VIAL 2 9305 9HCJ GCMS-2SRC C.. SAMPLES BATCHED SINGLE ANALYSIS
74 PLASTIC 1 7100 7RAH METALS-ICP C.. See Individual Elements Below
75 PLASTIC 1 7101 7RAH ALUMINUM C.. < 6.60854E-03 MOLAR
76 PLASTIC 1 7102 7RAH BORON C.. < 3.77882E-03 MOLAR
77 PLASTIC 1 7103 7 CADMIUM C.. NOT REQUESTED
78 PLASTIC 1 7105 7RAH IRON C.. < 2.22432E-04 MOLAR
79 PLASTIC 1 7110 7RAH ZIRCONIUM C.. < 1.57449E-03 MOLAR
80 PLASTIC 1 7111 7RAH CALCIUM C.. < 2.55539E-03 MOLAR
81 4 47981 7MLE SP-GR C.. 1.44216+-7.8057E-04.6.2574
                                                     784 CALCIUM C.. < 2.55539E-03 MOLAR
7981 7MLE SP-GR C.. 1.44216+-7.8057E-04 @ 25/4
7961 7MLE SPECTCHEM C.. Prep Completed 082393 14:53
7985 7CWL FLASHPOINT A.. Flashed @ 53.0 deg C corrected
7985 7CWL FLASHPOINT A.. NO Flash @ 60.0 deg C corrected
7985 7CWL FLASHPOINT A.. NO Flash @ 60.0 deg C corrected
7100 7GDD METALS-ICP C.. See Individual Elements Below
7101 7GDD ALUMINUM A.. 1.41894+-3.90244E-02 MOLAR
7981 7GDD SP-GR A.. 1.22883+-7.80484E-04 @ 25/4
7981 7GDD SP-GR A.. 1.22682+-7.80483E-04 @ 25/4
 81 4
 163 PLASTIC 2
 164 BEG KNOWN
165 RAMP 100
166 60C 100
 175 PLASTIC 2
176 PLASTIC 2
179 20
180 6
                                                               47981 7GDD SP-GR A. 1.22682+-7.80483E-04 @ 25/4
47981 7GDD SP-GR A. 1.24472+-7.80487E-04 @ 25/4
47981 7GDD SP-GR A. 1.24662+-7.80486E-04 @ 25/4
181 11
                                                    47981 7GDD SP-GR A.. 1.24662+-7.80486E-04 @ 25/4
7100 7WDT METALS-ICP C.. See Individual Elements Below
7105 7WDT IRON A.. 5.12682E-03+-3.06266E-04 MOLAR
77074 7MLE NITRATE A.. 6.20227+-.484159 Molar
7900 7VJJ EPA-TOX C.. See Individual Elements Below
7907 7VJJ SILVER A.. < 2.10773 mg/L
7900 7VJJ EPA-TOX C.. See Individual Elements Below
7907 7VJJ SILVER A.. < 2.10773 mg/L
7907 7VJJ SILVER A.. < 2.10773 mg/L
7908 7MLE SULFATE A.. 2.84473E-03+-2.36248E-04 Molar
63993 3DAO CS-134 A.. 12600+-714 D/S/ML
63993 3DAO CS-137 A.. 677000+-21000 D/S/ML
7961 7MLE AS/SE C.. Prep Completed 090993 04:05
7961 7MLE AS/SE C.. Prep Completed 090993 04:05
7961 7MLE AS/SE C.. Prep Completed 090993 04:06
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN ARSENIC A.. < 3.89309 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN ARSENIC A.. < 3.89309 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7901 7SDN SELENIUM A.. < .991639 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7906 7SDN SELENIUM A.. < .991639 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7906 7SDN SELENIUM A.. < .991639 mg/L
7900 7SDN EPA-TOX C.. See Individual Elements Below
7906 7SDN SELENIUM C.. < .991639 mg/L
7900 7CBG METALS-ICP C.. See Individual Elements Below
7906 7SDN SELENIUM C.. < .991639 mg/L
7100 7CBG METALS-ICP C.. See Individual Elements Below
7906 7SDN SELENIUM C.. < .991639 mg/L
7100 7CBG METALS-ICP C.. See Individual Elements Below
7101 7CBG ALUMINUM C.. > .956797 MOLAR
183 PLASTIC1R
                                                                  7100 7WDT METALS-ICP C.. See Individual Elements Below
184 PLASTIC1R
185 3R
186 #11
187 #11
188 #20
189 #20
190 3R
191 5
192 5
198 PLASTIC 1
199 PLASTIC 2
207 PLASTIC 1R
208 PLASTIC 1R
209 PLASTIC 2R
210 PLASTIC 2R
211 PLASTIC 1R
212 PLASTIC 1R
213 PLASTIC 2R
214 PLASTIC 2R
215 PLASTIC 1R
216 PLASTIC 1R
217 1R
218 1R
                                                                  7102 7CBG BORON C.. < 4.07588E-02 MOLAR
7105 7CBG IRON C.. 2.37342E-02+-1.54434
219 1R
220 1R
                                                                                                                                                C.. 2.37342E-02+-1.54434E-03 MOLAR
                                                          7110 7CBG IRON C.. 2.3/342E-02+-1.34434E-03 MOLAR
7111 7CBG CALCIUM C.. < 8.16728E-03 MOLAR
7111 7CBG CALCIUM C.. < 1.27546E-02 MOLAR
7100 7CBG METALS-ICP C.. See Individual Elements Below
7101 7CBG ALUMINUM C.. > .039455 MOLAR
221 1R
222 1R
223 1R
224 1R
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Log #
              Request Name Log Type
                                                        Charge Num Log Approval Info
   I
   d Lab
                           Meth a Ana-
x ID
                                   b lyst Analyte ARL
                                                                     Result
 930822-4
            WM-100
                                                          13120-200-001940210 23:01 SPLMGT SPLMGT
                         13120-200-001940210 23:01 SPI
47981 7BKH SP-GR A. .998247+-7.79139E-04 @ 25/4
47981 7BKH SP-GR A. .99821+-7.79138E-04 @ 25/4
47981 7BKH SP-GR A. .998321+-7.79139E-04 @ 25/4
67015 7MLE ACID A. < 8.97114E-02 NACID
67015 7MLE ACID A. < 8.97114E-02 NACID
67920 7BET URANIUM A. < 1.16045E-03 G/L
7972 7RLC UDS A. NO VISIBLE SOLIDS
57171 7MLE CHLORIDE A. < 15.0301 UG/ML
82 1
83 2
84 3
 85 2
 86 3
 87 2
 89 2
                         77074 7SRT NITRATE A.. 5.44794E-05+-9.6136E-06 Molar
                         67093 7MLE FLUORIDE A.. < 1.33105E-03 MOLAR
                         77168 7BKH SULFATE A.. 1.85622E-06+-4.93205E-07 Molar
92 3
                         7961 7WDT RADIOCHEM C.. Prep Completed 082393 11:27
93 5
                         7961 7WDT SPEC-CHEM C.. Prep Completed 082393 11:28
94 PLASTIC 1
95 PLASTIC 2
                    7961 7WDT SPEC-CHEM C.. Prep Completed 082393 11:28
```

Log I	#	Reques	t Name	Lo:	g Type		Charge Num Log Approval Info
đ	Lab		Meth	a An	a <b>-</b>		
x			#		st Analyte	ARL	Result
96	5		63993	3DAO	GAMMA SCAN	C	** See Index 193. **
97	5		23381		TOTAL SR		<< 1765.77+-619.312 D/S/ML
98	5		3205	3IDG	TRANS-UA		< .137597 D/S/ML
99	5		23011	3JSJ	TRITIUM		< 5.6679 D/SEC/ML
100	PLASTIC	1	7900	7CBG	EPA-TOX		See Individual Elements Below
101	PLASTIC	1	7901	7CBG	ARSENIC		< .67931 mg/L
102	PLASTIC	1	7902	7CBG	BARIUM		< .875559 mg/L
103	PLASTIC	1	7903	7CBG	CADMIUM		< .863441 mg/L
104	PLASTIC	1	7904		CHROMIUM		< .830681 mg/L
105	PLASTIC	1	7905	7CBG	LEAD		< .573449 mg/L
	PLASTIC		7906	7CBG	SELENIUM	Α	.849554+309505 mg/L
107	PLASTIC	1	7907	7CBG	SILVER	Α	< .774906 mg/L
108	PLASTIC	1	7908	7CBG	NICKEL	Α	< .806555 mg/L
109	PLASTIC	2	7900	7	EPA-TOX	С.,	NOT REQUESTED
	PLASTIC		7901	7	ARSENIC	С	NOT REQUESTED
111	PLASTIC	2	7902	7	BARIUM	С	NOT REQUESTED
112	PLASTIC	2	7903	7	CADMIUM	С	NOT REQUESTED
113	PLASTIC	2	7904	7	CHROMIUM	С.,	NOT REQUESTED
114	PLASTIC	2	7905	7	LEAD	С	NOT REQUESTED
115	PLASTIC	2	7906	7	SELENIUM	С.,	NOT REQUESTED
116	PLASTIC	2	7907	7	SILVER	С	NOT REQUESTED
117	PLASTIC	2	7908	7	NICKEL	С.,	NOT REQUESTED
	PLASTIC		2809	2LBZ	MERCURY	Α	=Not Detected:DL= 0.0045 mg/l
119	PLASTIC	2	2809	2LBZ	MERCURY	С	= Insufficient sample
120	PLASTIC	1	2330	2	ARSENIC.	С.,	SEE METHOD 7901
121	PLASTIC	2	2330	2	ARSENIC.	С.,	
122	PLASTIC	1	2340	2	SELENIUM.	С.,	SEE METHOD 7906
	PLASTIC		2340	2	SELENIUM.	С	SEE METHOD 7906
124	PLASTIC	1	2110	2JSL	SODIUM	Α	=6.7E-05 Molarity
	PLASTIC	1	2190		POTASSIUM	Α	<b>_</b>
126			67016	7KFM	PH	Α	<b>-</b>
127			7016	7	PH	С.,	
	PLASTIC		7961		SPEC-CHEM		
	VOA VIAL		9305		GCMS-2SRC	С	
	VOA VIAL		9305		GCMS-2SRC	C	
	PLASTIC		7100				See Individual Elements Below
	PLASTIC		7101		ALUMINUM		< 6.56896E-03 MOLAR
	PLASTIC		7102		BORON	Α	
	PLASTIC						NOT REQUESTED
	PLASTIC		7105				< 2.211E-04 MOLAR
	PLASTIC		7110				< 1.56506E-03 MOLAR
	PLASTIC	1 .	7111		CALCIUM		< 2.54008E-03 MOLAR
138		•	47981		SP-GR		.99835+-7.79139E-04 @ 25/4
	PLASTIC	<b>Z</b>	7961				Prep Completed 082393 11:29
	KNOWN		7985				Flashed @ 41.0 deg C corrected
168			7985				NO Flash @ 60.0 deg C corrected
193	כ		63993	3DAO	CS-137	Α	.792+0595 D/S/ML

Log	#	Request Name	Log Type	Charge Num Log Approval Info
I a	T - 1-	<b>36</b> 13	L	
	Lab ID		a Ana-	
x	TD	#	b lyst Analyte	ARL Result
140		7961	7MLE RADIOCHEM	C Prep Completed 082393 21:34
141	1	63993	3DAO GAMMA SCAN	A ** See Index 194 thru 197. **
142	1	23381	3BJS TOTAL SR	A 3.91922+106201 UCI/ML
143	1.	2110	2JSL SODIUM	A =0.048 Molarity
144	1	2190	2JSL POTASSIUM	A =0.0050 Molarity
145	1	77168	7BKH SULFATE	A 7.53216E-04+-1.6743E-04 Molar
146	1	77074	7SRT NITRATE	A 6.74799+614459 Molar
147	1	57171	7MLE CHLORIDE	A 70.1416+-16.2677 UG/ML
148	1	7972	7RLC UDS	A * 11.2667 MG/ML
149	1	47981	7WDT SP-GR	A 1.21087+-7.80464E-04 @ 25/4
150	1	67015	7MLE ACID	A < 8.97114E-02 NACID
151	1	67093	7MLE FLUORIDE	A < 6.65439E-03 MOLAR
152	1	7961	7MLE SPECTCHEM	
153	1	67920	7MLE URANIUM	A 4.20262E-03+-8.93985E-04 G/L
154	1	7100	7SDN METALS-ICP	C See Individual Elements Below
155	1	7101	7SDN ALUMINUM	C > .190546 MOLAR
156	1	7102	7SDN BORON	A < 3.75619E-03 MOLAR
157	1	7103	7 CADMIUM	C NOT REQUESTED
158	1	7105	7SDN IRON	A 6.55824E-03+-3.42343E-04 MOLAR
159		7110	7SDN ZIRCONIUM	
160	1	7111	7SDN CALCIUM	A < 2.54008E-03 MOLAR
173		7100	7RAH METALS-ICP	C See Individual Elements Below
174	1	7101	7RAH ALUMINUM	A 1.39981+366096 MOLAR
177	1	7900	7WDT EPA-TOX	C See Individual Elements Below
178	1	7903	7WDT CADMIUM	A 11.9089+-1.07906 mg/L
194		63993	3DAO CO-60	A 129+-44.1 D/S/ML
195	1	63993	3DAO CS-134	A 10400+-376 D/S/ML
196	1	63993	3DAO CS-137	A 309000+-9770 D/S/ML
197	1	63993	3DAO EU-154	A 662+-89.7 D/S/ML
9309	01-23	WM-100AT		13120-200-001940204 13:14 SPLMGT SPLMGT
263	1	7961	7MLE SPECTROCH	
264	1	33993	3DAO GAMMA SCAN	C ** See Index 294 thru 295. **
265	1	23381	3JSJ TOTAL SR	A 4.25367E-03+-1.63083E-04 UCI/M
266	1	2110	2JSL SODIUM	A =0.050 Molarity
267	1	2190	2JSL POTASSIUM	A =5.4E-03 Molarity
268		77168		A 7.8067E-04+-8.24152E-05 Molar
269	1	77074	7MLE NITRATE	A 4.48163+645531 Molar
270		57171	7MLE CHLORIDE	C < 60.8519 UG/ML
271		7972	7MLE UDS	A * 2.33333 MG/ML
272		47981	7KFM SP-GR	A 1.24505+-7.80487E-04 @ 25/4
273		67015	7MLE ACID	A < 8.97114E-02 NACID
274		67093	7MLE FLUORIDE	A 7.14322E-03+-3.61877E-04 MOLAR
275		7961		C Prep Completed 090293 13:29
276		67920	7MLE URANIUM	A 3.59186E-03+-8.31949E-04 G/L
277 1		7100		C See Individual Elements Below
278		7101		C > .191694 MOLAR
279	L	7102	7WDT BORON	A < 3.77882E-03 MOLAR

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Log #
                           Request Name Log Type
                                                                                                  Charge Num Log Approval Info
    I
     d Lab
                                                 Meth a Ana-
     x ID
                                                                b lyst Analyte ARL
                                                                                                                                 Result
                                          7103 7WDT CADMIUM C.. < 9.5635E-04 MOLAR
7105 7WDT IRON A.. 2.38671E-02+-6.51053E-04 MOLAR
 280 1
 281 1
                                            7110 7WDT ZIRCONIUM A. < 1.57449E-03 MOLAR
7111 7WDT CALCIUM A. 4.29423E-03+-6.58732E-04 MOLAR
7961 7MLE RADIOCHEM C. Prep Completed 090293 13:29
7100 7WDT METALS-ICP C. See Individual Elements Below
7101 7WDT ALUMINUM A. 1.91275+-.040828 MOLAR
 282 1
 283 1
 284 1
 285 1
 286 1
                                         7701 7WD1 ABONINGS A.. 2.32.5.44 UG/ML
7900 7WDT EPA-TOX C.. See Individual Elements Below
7903 7WDT CADMIUM A.. 24.9983+-1.75358 mg/L
 287 1R
288 1
289 1
                                       57171 7KFM CHLORIDE A. 45.3354+-8.37305 UG/ML
7100 7WDT METALS-ICP C. See Individual Elements Below
7101 7WDT ALUMINUM A. 1.41061+-.147991 MOLAR
290 2R
291 PLASTIC1R
292 PLASTIC1R
294 1
                                            33993 3DAO CS-134 A.. 14700+-656 D/S/ML
295 1
                                            33993 3DAO CS-137
                                                                                                A.. 438000+-13800 D/S/ML
930903-19 WM-100AT
                                                                                                             13120-200-001930922 15:02 SPLMGT SPLMGT
728 1 7961 7BET RADIO CHEM C.. Prep Completed 090493 19:29
729 1
                                               63993 3DAO GAMMA SCAN C.. ** See Index 820 thru 823.
730 1
                                             23381 3BJS TOTAL SR A.. 4.08916+-.102223 UCI/ML
731 1
                                             2110 2JSL SODIUM A.. =0.049 Molarity
2190 2JSL POTASSIUM A.. =6.0E-03 Molarity
732 1
733 1
                                           77168 7BET SULFATE A.. 4.9744E-04+-7.58426E-05 Molar
77074 7BET NITRATE A.. 3.94383+-.23135 Molar
57171 7RLC CHLORIDE A.. 23.156+-4.18063 UG/ML
734 1
734 -
735 1
736 1
                                        77074 7BET NITRATE A.. 3.94383+-.23135 Molar
57171 7RLC CHLORIDE A.. 23.156+-4.18063 UG/ML
7972 7RAH UDS A.. * 1.68 MG/ML
47981 7RAH SP-GR A.. 1.23005+-7.80485E-04 @ 25/4
67015 7BGP ACID A.. < .136135 NACID
67093 7BGP FLUORIDE A.. < 7.96535E-03 MOLAR
7961 7BET SPECT CHEM C.. Prep Completed 090493 19:30
67920 7RLC URANIUM A.. 1.04237E-02+-1.37579E-03 G/L
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM C.. > .190546 MOLAR
7102 7SDN BORON A.. < 4.84314E-03 MOLAR
7103 7 CADMIUM C.. SEE INDEX 809
7105 7SDN IRON A.. 1.84785E-02+-6.4824E-04 MOLAR
7110 7SDN ZIRCONIUM A.. < 1.39699E-03 MOLAR
7111 7SDN CALCIUM A.. 4.1136E-03+-6.3767E-04 MOLAR
47981 7RAH SP-GR A.. 1.22971+-7.80485E-04 @ 25/4
47981 7RAH SP-GR A.. 1.22971+-7.80484E-04 @ 25/4
47981 7RAH SP-GR A.. 1.22997+-7.80484E-04 @ 25/4
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM A.. 1.6911+-7.25564E-02 MOLAR
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM A.. 1.6911+-7.25564E-02 MOLAR
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN CADMIUM A.. < 1.30499E-03 MOLAR
63993 3DAO CO-60 A.. 311+-54.5 D/S/ML
63993 3DAO CS-134 A.. 12100+-442 D/S/ML
63993 3DAO CS-137 A.. 371000+-10500 D/S/ML
737 1
738 1
739 1
740 1
741 1
742 1
743 1
744 1
745 1
746 1
747 1
748 1
791 2
792 3
793 4
804 1
805 1
808 1
809 1
820 1
821 1
822 1
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Log # I	Request Name	Log Type L	1	Charge Num Log Approval Info
d Lab	Meth	a Ana-		
x ID	#		ARL	Result
823 1	63993	3DAO EU-154	Α	768+-109 D/S/ML
930903-20	WM-100AT			13120-200-001930922 15:17 SPLMGT SPLMGT
749 1	7961	7BET RADIO CHEM	С	Prep Completed 090493 19:35
750 1	73993			** See Index 824 thru 827. **
751 1	23381	3BJS TOTAL SR		4.25436+104309 UCI/ML
752 1	2110	2JSL SODIUM	Α	=0.051 Molarity
753 1	2190	2JSL POTASSIUM	Α	=6.0E-03 Molarity
754 1	77168	7BET SULFATE		4.35997E-04+-7.22972E-05 Molar
755 1	77074	7BET NITRATE	Α	3.9671+23124 Molar
756 1	57171	7RLC CHLORIDE		17.0564+-4.02563 UG/ML
757 1	7972	7RAH UDS		* 1.19 MG/ML
758 1	47981	7RAH SP-GR		1.22274+-7.8048E-04 @ 25/4
759 1	67015	7BGP ACID		< .136135 NACID
760 1	67093	7BGP FLUORIDE		< 7.96535E-03 MOLAR
761 1	7961			Prep Completed 090493 19:30
762 1	67920	7RLC URANIUM		3.78881E-03+-8.52444E-04 G/L
763 1	7100			See Individual Elements Below
764 1	7101	7SDN ALUMINUM		> .190546 MOLAR
765 1	7102	7SDN BORON		< 4.84314E-03 MOLAR
766 1	7103	7 CADMIUM	С	
767 1	7105	7SDN IRON	Α	
768 1	7110			< 1.39699E-03 MOLAR
769 1 79 <b>4</b> 2	7111	7SDN CALCIUM		4.17561E-03+-6.40325E-04 MOLAR
794 2 795 3	47981	7RAH SP-GR		1.23732+-7.80487E-04 @ 25/4
796 4	47981	7RAH SP-GR	A	·
806 1	47981 7100	7RAH SP-GR	A	· · · · · · · · · · · · · · · · · · ·
807 1	7100	7SDN METALS-ICP 7SDN ALUMINUM	A	See Individual Elements Below 1.33428+-6.51375E-02 MOLAR
810 1	7101			See Individual Elements Below
811 1	7103	7SDN METALS-ICP 7SDN CADMIUM		< 1.30499E-03 MOLAR
824 1	73993	3DAO CO-60	A	·
825 1	73993			14600+-508 D/S/ML
826 1	73993	3DAO CS-134 3DAO CS-137		486000+-15100 D/S/ML
827 1	73993	3DAO EU-154		1170+-115 D/S/ML
-		22.10 20 201		22,0, 240 2,0,112
930903-21	WM-100AT			13120-200-001930922 15:16 SPLMGT SPLMGT
770 1	7961	7BET RADIO CHEM	С	Prep Completed 090493 19:36
771 1	83993			** See Index 828 thru 831. **
772 1	23381	3BJS TOTAL SR	Α	
773 1	2110	2JSL SODIUM	Α	<u>.</u>
774 1	2190	2JSL POTASSIUM		=5.9E-03 Molarity
775 1	77168	7BET SULFATE		NO3 SPL CTS HIGHER THAN C.S. CT
776 1	77074	7BET NITRATE	Α	4.4177+229134 Molar
777 1	57171	7RLC CHLORIDE		24.0692+-4.20335 UG/ML
778 1	7972	7RLC UDS	Α	* 1.55789 MG/ML
779 1	47981	7RAH SP-GR	Α	· · · · · · · · · · · · · · · · · · ·
780 1	67015			< .136135 NACID
781 1	67093	7BGP FLUORIDE	С	< 7.96535E-03 MOLAR

Log	#	Request Name	Log Type L	Charge Num Log Approval Info
đ	Lab	Meth	a Ana-	
x	ID	#	b lyst Analyte	ADI Damile
		π	b Tyst Analyte	ARL Result
782	1	7961	7BET SPECT CHEM	C Prep Completed 090493 19:31
783	1	67920	7RLC URANIUM	A 3.59557E-03+-8.3234E-04 G/L
784	1	7100	7SDN METALS-ICP	C See Individual Elements Below
785	1	7101	7SDN ALUMINUM	C > .190546 MOLAR
786	1	7102	7SDN BORON	A < 4.84314E-03 MOLAR
787	1	7103	7 CADMIUM	C SEE INDEX 813
788	1	7105	7SDN IRON	A018899+-6.55724E-04 MOLAR
789		7110	7SDN ZIRCONIUM	A < 1.39699E-03 MOLAR
790	1	7111	7SDN CALCIUM	A 4.33933E-03+-6.47285E-04 MOLAR
797	2	47981	7RAH SP-GR	A 1.23281+-7.80486E-04 @ 25/4
798	3	47981	7RAH SP-GR	A 1.22992+-7.80485E-04 @ 25/4
799	4	47981	7RAH SP-GR	A 1.22116+-7.80478E-04 @ 25/4
800	1	67093	7BGP FLUORIDE	A 9.76166E-03+-1.30172E-03 MOLAR
801	2	77168	7BET SULFATE	A 8.04588E-04+-1.07187E-04 Molar
802	1	7100	7SDN METALS-ICP	C See Individual Elements Below
803	1	7101	7SDN ALUMINUM	A 1.4426+-6.74725E-02 MOLAR
812	1	7100	7SDN METALS-ICP	C See Individual Elements Below
813	1	7103	7SDN CADMIUM	A < 1.30499E-03 MOLAR
828	1	83993	3DAO CO-60	A 470+-64.8 D/S/ML
829	1	83993	3DAO CS-134	A 14000+-504 D/S/ML
830	1	83993	3DAO CS-137	A 438000+-14300 D/S/ML
831	1	83993	3DAO EU-154	A 1190+-140 D/S/ML

\*\*\*\*\*\*\* END \*\*\*\*\*\*

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Date of Search: 2003-06-09 16:27:03.545
                                                  Run by: JEFF LONG
**************************
Search Criteria:
   Start Log....:930101 1
   End Log.....:931231 14
   Log Approval .: ALL Logs
   Result Type ... : All Entries
   Lab/Group....:ALL Groups
   Name Column..:Lab Sample ID
    Request Name..:*WM-102*
Total # Logs Found...: 6
Total # Results Found: 432
**********
Log #
            Request Name Log Type
                                          Charge Num
                                                         Log Approval Info
  I
  d Lab
                     Meth a Ana-
  x ID
                     # b lyst Analyte ARL
                                                     Result
930505-14 WM-102:CH
                                            13120-200-001940209 23:08 SPLMGT SPLMGT
                                      A.. 1.17066+-6.89039E-04 @ 25/4
508 1
                   47981 7MLE SP-GR
509 2
                   47981 7MLE SP-GR
                                       A.. 1.17051+-6.8904E-04 @ 25/4
510 3
                   47981 7MLE SP-GR
                                       A.. 1.17048+-6.8904E-04 @ 25/4
511 4
                  47981 7MLE SP-GR
                                       A.. 1.17053+-6.8904E-04 @ 25/4
                  67015 7MLE ACID A.. < .229527 NACID
67015 7MLE ACID A.. < .229527 NACID
512 2
513 3
                  67920 7BGP URANIUM A.. < 4.25198E-03 G/L
514 2
                   7972 7SDN UDS A.. * 352.041 UG/ML
515 4
516 2
                  57171 7MLE CHLORIDE A.. < 36.3961 UG/ML
517 3
                  77074 7SLI NITRATE A.. 2.70904+-.178567 Molar
518 2
                  67093 7AWO FLUORIDE A.. 90.8023+-12.1744 UG/ML
519 3
                  77168 7SLI SULFATE C.. < 2.76176E-02 Molar
520 5
                   7961 7MLE RADIOCHEM C.. Prep Completed 050693 05:26
521 PLASTIC 1
                   7961 7MLE SPEC CHEM C.. Prep Completed 050693 05:27
522 PLASTIC 2
                   7961 7MLE SPEC CHEM C.. Prep Completed 050693 05:28
                  33993 3DAO GAMMA SCAN C.. ** See Index 812 thru 814.
523 5
524 5
                 23381 3BJS TOTAL SR A.. 37094.9+-1409.7 D/S/ML
525 5
                  3998 3 TRANS-UA C.. SEE INDEX 738 AND 739
526 5
                 23011 3JSJ TRITIUM A.. < 16.3201 D/SEC/ML
527 PLASTIC 1
                7100 7SDN METALS-ICP C.. See Individual Elements Below
528 PLASTIC 1
                   7101 7SDN ALUMINUM C.. > .186818 MOLAR
                  7102 7SDN BORON A.. < 3.68268E-03 MOLAR
7105 7SDN IRON A.. 2.88252E-02+-7.08747E-04 MOLAR
529 PLASTIC 1
530 PLASTIC 1
531 PLASTIC 1
                  7110 7SDN ZIRCONIUM A.. < 1.53443E-03 MOLAR
                 7111 7SDN CALCIUM A.. < 2.49037E-03 MOLAR
7900 7RAH EPA-TOX C.. See Individual Elements Below
7901 7RAH ARSENIC C.. < 29.4377 mg/L
7902 7RAH BARIUM A.. < 32.6799 mg/L
532 PLASTIC 1
533 PLASTIC 1
534 PLASTIC 1
535 PLASTIC 1
               7903 7RAH CADMIUM C.. < 42.3012 mg/L
```

536 PLASTIC 1

Log I	#	Request	Name	Log	g Type		Charge Num Log Approval Info
đ	Lab		Meth	a Ana	a-		
x	ID				st Analyte	ARL	Result
	PLASTIC		7904		CHROMIUM		< 30.983 mg/L
	PLASTIC		7905		LEAD		< 43.9272  mg/L
	PLASTIC		7906		SELENIUM		49.1119+-14.5269 mg/L
	PLASTIC		7907		SILVER		< 26.5997 mg/L
	PLASTIC		7908		NICKEL		< 47.2862 mg/L
	PLASTIC		7900	7RAH	EPA-TOX		See Individual Elements Below
	PLASTIC		7901	7RAH	ARSENIC		< 29.4377 mg/L
	PLASTIC		7902		BARIUM		< 32.6799 mg/L
	PLASTIC		7903		CADMIUM		< 42.3012 mg/L
	PLASTIC		7904		CHROMIUM		< 30.983 mg/L
	PLASTIC		7905	7RAH	LEAD		< 43.9272 mg/L
	PLASTIC		7906		SELENIUM		37.0847+-13.0129 mg/L
	PLASTIC		7907		SILVER		< 26.5997 mg/L
	PLASTIC		7908	7RAH	NICKEL	Α	< 47.2862  mg/L
	PLASTIC		2809		MERCURY	Α	=Not Detected:DL= 0.075 mg/l
	PLASTIC		2809	2LBZ	MERCURY	Α	
	PLASTIC		2330	2	ARSENIC.		SEE RAL DATA
	PLASTIC		2330	2	ARSENIC.		SEE RAL DATA
	PLASTIC		2340	2	SELENIUM.	С.,	SEE RAL DATA SEE RAL DATA
	PLASTIC		2340	2	SELENIUM.	С	SEE RAL DATA
	PLASTIC		2119	2JSL	SODIUM	Α	=1.0E-02 Molarity
	PLASTIC	1	2199	2JSL	POTASSIUM	Α	=8.2E-03 Molarity
559			57016	7AWO	PH	Α	< .5 pH
	PLASTIC	1	7961				Prep Completed 050693 05:29
	5 RAMP			7CWL	FLASHPOINT	С	NO Flash @ 60.0 deg C corrected
	BEG KNW		7985	7CWL	FLASHPOINT	С	Flashed @ 60.0 deg C corrected
	VOA VIAL		9304		GCMS-2TRAP		· · · · · · · · · · · · · · · · · · ·
	VOA VIAL	. 2	9304			Α	SEE ATTACHED
738			3202		PU-238	Α	
739			3203		PU-239		< 74.2257 D/S/ML
751			7900				See Individual Elements Below
752			7903		CADMIUM		9.49254 + -1.37944  mg/L
753			7904		CHROMIUM		9.35827+-1.83895 mg/L
754			7905		LEAD		< 4.13354  mg/L
755			7907		SILVER		< 2.50302 mg/L
756			7900				See Individual Elements Below
757			7903		CADMIUM		30.5871+-2.73251 mg/L
758			7904		CHROMIUM		32.1589+-3.17676 mg/L
759			7905		LEAD		< 4.13354 mg/L
760		_	7907		SILVER		< 2.50302 mg/L
795					SULFATE		1.27844E-03+-8.11336E-05 Molar
812					RU-106		9250+-1050 D/S/ML
813					CS-134		35800+-1360 D/S/ML
814					CS-137	Α	• • • • • • • • • • • • • • • • • • • •
	PLASTIC						See Individual Elements Below
	PLASTIC	Т	7101	_	ALUMINUM	A	
	5 RAMP		7985	7	FLASHPOINT		
	60 CHK						NO Flash @ 60.0 deg C corrected
845	PLASTIC	Τ	7900	/WDT	EPA-TOX	С	See Individual Elements Below

```
Log #
              Request Name
                            Log Type
                                               Charge Num Log Approval Info
  I
                             Τ.
  d Lab
                       Meth a Ana-
  x ID
                             b lyst Analyte ARL
                                                           Result
846 PLASTIC 1
                      7903
                             7WDT CADMIUM
                                              C.. 45.479+-2.39422 mg/L
847 PLASTIC 1
                      7904
                             7WDT CHROMIUM
                                              C.. 45.9642+-3.9403 mg/L
                   7904 /WDT CHROMIUM C.. 45.5042+-3.5403 mg/L
7900 7 EPA-TOX C.. NOT REQUESTED
7900 7WDT EPA-TOX C.. See Individual Elements Below
7906 7WDT SELENIUM C.. < 2.56205 mg/L
7900 7WDT EPA-TOX C.. See Individual Elements Below
7906 7WDT SELENIUM C.. < 2.56205 mg/L
7906 7WDT SELENIUM C.. < 2.56205 mg/L
7907 7RAH AS/SE PREP C.. Prep Completed 091093 10:19
7908 7RAH SPECTROCHE C.. Prep Completed 091093 10:19
848 PLASTIC 1
867 PLASTIC 1
                                             C.. See Individual Elements Below
868 PLASTIC 1
869 PLASTIC 2
                                             C.. See Individual Elements Below
870 PLASTIC 2
876 PLASTIC 1
877 PLASTIC 2
883 PLASTIC 1R
                     7900 7JMK EPA-TOX
                                           C.. See Individual Elements Below
884 PLASTIC 1R
                     7901 7JMK ARSENIC
                                             A... < 3.89309 \text{ mg/L}
885 PLASTIC 2R
                     7900 7JMK EPA-TOX
                                             C.. See Individual Elements Below
886 PLASTIC 2R
                     7901 7JMK ARSENIC
                                           A.. < 3.89309 mg/L
887 PLASTIC 1R
                     7900 7JMK EPA-TOX
                                             C.. See Individual Elements Below
888 PLASTIC 1R
                     7906 7JMK SELENIUM A.. < .991639 mg/L
889 PLASTIC 2R
                     7900 7JMK EPA-TOX
                                             C.. See Individual Elements Below
                      7906 7JMK SELENIUM A.. < .991639 mg/L
890 PLASTIC 2R
                      7900 7SDN EPA-TOX
911 PLASTIC 1R
                                             C.. See Individual Elements Below
912 PLASTIC 1R
                      7903 7SDN CADMIUM
                                             A.. 31.3152+-1.52779 mg/L
913 PLASTIC 1R
                      7904 7SDN CHROMIUM A.. 30.1857+-3.00239 mg/L
916 SPIKE
                      9304 9HCJ GCMS-2TRAP A.. SEE ATTACHED
930505-15 WM-102:CH
                                                  13120-200-001940209 22:32 SPLMGT SPLMGT
                     47981 7MLE SP-GR
                                             A.. 1.16163+-6.89136E-04 @ 25/4
566 2
                     47981
                            7MLE SP-GR
                                             A.. 1.17057+-6.89039E-04 @ 25/4
567 3
                     47981
                            7MLE SP-GR
                                             A.. 1.17056+-6.89039E-04 @ 25/4
                            7MLE SP-GR
568 4
                                            A.. 1.17052+-6.8904E-04 @ 25/4
                     47981
569 2
                    67015 7MLE ACID
                                            A.. < .229527 NACID
570 3
                    67015 7MLE ACID
                                            A.. < .229527 NACID
571 2
                   67920 7BGP URANIUM A.. < 4.25198E-03 G/L
                                            A.. * 1755.32 UG/ML
572 4
                     7972 7SDN UDS
573 2
                    57171 7MLE CHLORIDE A.. < 36.3961 UG/ML
                                             A.. 2.77898+-.178497 Molar
574 3
                    77074 7SLI NITRATE
575 2
                   67093 7AWO FLUORIDE A.. 86.2272+-11.9815 UG/ML
576 3
                    77168 7SLI SULFATE
                                             C.. < 2.76176E-02 Molar
577 5
                     7961 7MLE RADID CHEM C.. Prep Completed 050693 05:29
                     7961 7MLE RADID CHEM C.. Prep Completed 050693 05:30
578 PLASTIC 1
                     7961 7MLE RADID CHEM C.. Prep Completed 050693 05:30
579 PLASTIC 2
                    33993 3DAO GAMMA SCAN C.. ** See Index 815 thru 816.
580 5
581 5
                   23381 3BJS TOTAL SR A.. 37352.3+-1414.36 D/S/ML
582 5
                     3998 3
                                  TRANS-UA C.. SEE INDEX 740 AND 741
                   23011
583 5
                            3JSJ TRITIUM
                                            A.. < 16.3201 D/SEC/ML
                    7100
584 PLASTIC 1
                            7SDN METALS-ICP C.. See Individual Elements Below
585 PLASTIC 1
                     7101
                            7SDN ALUMINUM C.. > .189789 MOLAR
586 PLASTIC 1
                     7102 7SDN BORON
                                         A.. < 3.74125E-03 MOLAR
587 PLASTIC 1
                     7105
                            7SDN IRON
                                            A.. 3.31979E-02+-7.68484E-04 MOLAR
588 PLASTIC 1
                    7110
                            7SDN ZIRCONIUM A.. < 1.55883E-03 MOLAR
589 PLASTIC 1
                    7111
                            7SDN CALCIUM A.. < 2.52998E-03 MOLAR
                    7900
590 PLASTIC 1
                            7RAH EPA-TOX
                                             C.. See Individual Elements Below
```

Log I	#	Request	Name	Lo.	g Type		Charge Num Log Approval Info
đ	Lab		Meth		a-		
<b>x</b>	ID		#		st Analyte	ARL	Result
	PLASTIC	_	7901	7RAH	ARSENIC	С	< 29.4377 mg/L
	PLASTIC		7902	7RAH	BARIUM	Α	< 32.6799 mg/L
	PLASTIC		7903		CADMIUM		< 42.3012 mg/L
	PLASTIC		7904	7RAH	CHROMIUM	С.,	< 30.983 mg/L
	PLASTIC		7905	7RAH	LEAD	С	<u> </u>
	PLASTIC		7906		SELENIUM		< 24.4469 mg/L
	PLASTIC		7907	7RAH	SILVER		< 26.5997 mg/L
	PLASTIC	_	7908		NICKEL		< 47.2862 mg/L
	PLASTIC		7900		EPA-TOX		See Individual Elements Below
	PLASTIC		7901		ARSENIC		< 29.4377 mg/L
	PLASTIC		7902		BARIUM	Α	
	PLASTIC		7903		CADMIUM		< 42.3012 mg/L
	PLASTIC		7904		CHROMIUM		< 30.983 mg/L
	PLASTIC		7905		LEAD		< 43.9272 mg/L
	PLASTIC		7906		SELENIUM		32.7313+-12.4197 mg/L
	PLASTIC		7907			С	
	PLASTIC		7908		NICKEL		< 47.2862 mg/L
	PLASTIC		2809		MERCURY	Α	
	PLASTIC		2809		MERCURY	Α	
	PLASTIC		2330	2	ARSENIC.	C	
	PLASTIC		2330	2	ARSENIC.	C	SEE RAL DATA SEE RAL DATA
	PLASTIC		2340	2	SELENIUM.		
	PLASTIC		2340	2	SELENIUM.		SEE RAL DATA
	PLASTIC		2119		SODIUM	Α	<del>-</del>
616	PLASTIC		2199		POTASSIUM		<del>-</del>
			7016	7AWO			< .5 pH
	PLASTIC 5 RAMP	1	7961	/MLE	RADID CHEM	C	Prep Completed 050693 05:30
	END KNOW	NT	7985 7985				NO Flash @ 60.0 deg C corrected
	VOA VIAL						Flashed @ 60.0 deg C corrected
	VOA VIAL		9304 9304		GCMS-2TRAP		
740		2	3202		PU-238	A	
741			3202		PU-239		2878.94+-101.325 D/S/ML < 74.2257 D/S/ML
761			7900		EPA-TOX		See Individual Elements Below
762			7903		CADMIUM	A	
763			7904		CHROMIUM	Α	3.
764			7905		LEAD		< 4.13354 mg/L
765			7907		SILVER	A	
766			7900		EPA-TOX		See Individual Elements Below
767			7903		CADMIUM		21.9886+-2.27864 mg/L
768			7904		CHROMIUM		21.7697+-2.64986 mg/L
769			7905		LEAD		< 4.13354 mg/L
770			7907		SILVER	A	<b>5</b> .
796		7	7168		SULFATE		1.35747E-03+-8.25489E-05 Molar
815			3993		CS-134		41900+-1820 D/S/ML
816					CS-137		559000+-24900 D/S/ML
	PLASTIC :		7100				See Individual Elements Below
	PLASTIC :		7101		ALUMINUM	Α	
	60 CHK		7985				NO Flash @ 60.0 deg C corrected
							-

```
Log #
                                       Request Name Log Type
                                                                                                                                      Charge Num Log Approval Info
     I
        d Lab
                                                                    Meth a Ana-
        x ID
                                                                                         b lyst Analyte ARL
                                                                                                                                                                                      Result

      849 VOA VIAL
      7900 7WDT EPA-TOX
      C... See Individual Elements Below

      850 VOA VIAL
      7903 7WDT CADMIUM
      C... 11.9906+-1.07472 mg/L

      851 VOA VIAL
      7904 7WDT CHROMIUM
      C... 5.93173+-1.14071 mg/L

      852 VOA VIAL
      7906 7WDT SELENIUM
      C... < 2.56205 mg/L</td>

      853 VOA VIAL
      7900 7WDT EPA-TOX
      C.. See Individual Elements Below

      854 VOA VIAL
      7906 7WDT SELENIUM
      C... < 2.56205 mg/L</td>

      878 PLASTIC 1
      7961 7KFM SP CHEM
      C... Prep Completed 090993 20:49

      879 PLASTIC 1
      7961 7KFM SP CHEM
      C... Prep Completed 090993 20:50

      880 PLASTIC 2
      7961 7KFM SP CHEM
      C... Prep Completed 090993 20:50

      891 PLASTIC 1R
      7900 7JMK EPA-TOX
      C.. See Individual Elements Below

      892 PLASTIC 1R
      7901 7JMK ARSENIC
      A.. < 3.89309 mg/L</td>

      893 PLASTIC 2R
      7901 7JMK ARSENIC
      A.. < 3.89309 mg/L</td>

      894 PLASTIC 1R
      7900 7JMK EPA-TOX
      C.. See Individual Elements Below

      896 PLASTIC 1R
      7900 7JMK SELENIUM
      A.. < 3.89309 mg/L</td>

      897 PLASTIC 2R
      7906 7JMK SELENIUM
      A.. < 991639 mg/L</td>

      898 PLASTIC 2R
      7906 7JMK SELENIUM
      A.. < .991639 mg/L</td>

      898 PLASTIC 2R
      7906 7JMK SELENIUM
  849 VOA VIAL 7900
                                                                                        7WDT EPA-TOX
                                                                                                                                             C.. See Individual Elements Below
                                                                   9304 9HCJ GCMS-2TRAP A.. SEE ATTACHED
  917 SPIKE
  930505-16 WM-102:CH
                                                                                                                                                        13120-200-001940221 10:25 SPLMGT SPLMGT
                                                                47981 7MLE SP-GR A. 1.16789+-6.89067E-04 @ 25/4
47981 7MLE SP-GR A. 1.17065+-6.89039E-04 @ 25/4
47981 7MLE SP-GR A. 1.17053+-6.8904E-04 @ 25/4
47981 7MLE SP-GR A. 1.17055+-6.8904E-04 @ 25/4
  622 1
  623 2
  624 3
  625 4
 626 2
                                                               67015 7MLE ACID A.. < .229527 NACID
67015 7MLE ACID A.. < .229527 NACID
 627 3
                                                               67015 7MLE ACID
 628 2
                                                             67920 7BGP URANIUM A.. < 4.25198E-03 G/L
                                                                 7972 7SDN UDS A.. * 8855.55 UG/ML
57171 7MLE CHLORIDE A.. < 36.3961 UG/ML
 629 4
 630 2
                                                                 77074
 631 3
                                                                                       7SLI NITRATE A.. 2.69894+-.178578 Molar
                                                             77074 75H NITRATE A. 2.098947-176378 MOTAL
67093 7AWO FLUORIDE A. 58.2337+-10.7303 UG/ML
77168 7SLI SULFATE A. 2.88159E-02+-9.03836E-03 Molar
7961 7MLE RADIO CHEM C. Prep Completed 050693 05:30
7961 7MLE SPEC CHEM C. Prep Completed 050693 05:31
7961 7MLE SPEC CHEM C. Prep Completed 050693 05:33
33993 3DAO GAMMA SCAN C. ** See Index 817 thru 821.
 632 2
 633 3
 634 5
 635 PLASTIC 1
 636 PLASTIC 2
                                                    7961 /MLE SPEC CHEAR C.. 122
33993 3DAO GAMMA SCAN C.. ** See Index 817 thru 821.
23381 3BJS TOTAL SR A.. 37270.1+-1412.87 D/S/ML
3998 3 TRANS-UA C.. SEE INDEX 742 AND 743
23011 3JSJ TRITIUM A.. < 16.3201 D/SEC/ML
7100 7SDN METALS-ICP C.. See Individual Elements Below
7101 7SDN ALUMINUM C.. < 8.65524E-03 MOLAR
7102 7SDN BORON A.. < 3.68268E-03 MOLAR
7105 7SDN IRON A.. < 2.16774E-04 MOLAR
 637 5
 638 5
 639 5
 640 5
 641 PLASTIC 1
 642 PLASTIC 1
 643 PLASTIC 1
 644 PLASTIC 1
```

Log I	#	Request	. Name	Lo.	g Type		Charge Num Log Approval Info
đ	Lab		Meth	a An	a-		
х	ID		#	b ly	st Analyte	ARL	Result
645	PLASTIC	1	7110	7SDN	ZIRCONIUM	Α	< 1.53443E-03 MOLAR
646	PLASTIC	1	7111	7SDN	CALCIUM	Α	< 2.49037E-03 MOLAR
647	PLASTIC	1	7900	7RAH	EPA-TOX	С	See Individual Elements Below
	PLASTIC		7901	7RAH	ARSENIC	С	< 29.4377 mg/L
649	PLASTIC	1	7902	7RAH	BARIUM	С.,	< 32.6799 mg/L
650	PLASTIC	1	7903	7RAH	CADMIUM		< 42.3012 mg/L
	PLASTIC		7904	7RAH	CHROMIUM		< 30.983 mg/L
	PLASTIC		7905	7RAH	LEAD		< 43.9272 mg/L
653	PLASTIC	1	7906	7RAH	SELENIUM	С	29.0911+-11.9012 mg/L
	PLASTIC		7907	7RAH	SILVER	С	< 26.5997 mg/L
	PLASTIC	1	7908	7RAH	NICKEL		< 47.2862 mg/L
656			7900		EPA-TOX	С.,	See Individual Elements Below
657			7901	7WDT	ARSENIC	Α	< 2.77008 mg/L
658			7902		BARIUM		< 3.07517 mg/L
659			7903	7WDT	CADMIUM	Α	29.6708+-2.68771 mg/L
660			7904	7WDT	CHROMIUM	Α	32.1341+-3.17561  mg/L
661			7905		LEAD		< 4.13354 mg/L
662			7906	7wdr	SELENIUM		< 2.30045 mg/L
663			7907		SILVER		< 2.50302 mg/L
664			7908		NICKEL	Α	5.5652+375134 mg/L
	PLASTIC		2809		MERCURY	Α	
	PLASTIC		2809	2LBZ	MERCURY	Α	•
	PLASTIC		2330	2	ARSENIC.	С.,	
	PLASTIC		2330	2	ARSENIC.	С	
	PLASTIC		2340	2	SELENIUM.		SEE RAL DATA
	PLASTIC		2340	2	SELENIUM.	С.,	SEE RAL DATA
	PLASTIC		2119		SODIUM		=9.5E-03 Molarity
	PLASTIC	1	2199		POTASSIUM	Α	<u>-</u>
673			67016	7SDN			< .5 pH
	PLASTIC	1	7961				Prep Completed 050693 05:33
	5 RAMP	<b></b>					NO Flash @ 60.0 deg C corrected
	BEG KNOW		7985				Flashed @ 60.0 deg C corrected
	VOA VIAL		9304		GCMS-2TRAP		
742	VOA VIAL	ı 4	9304		GCMS-2TRAP		
743			3202 3203		PU-238 PU-239	A	• •
771							< 74.2257 D/S/ML
772			7900		EPA-TOX		See Individual Elements Below
773			7903 7904		CADMIUM CHROMIUM		29.201+-2.66446 mg/L 30.9679+-3.1207 mg/L
774			7904		LEAD		< 4.13354 mg/L
775			7905		SILVER		< 4.13354 mg/L < 2.50302 mg/L
817			33993		CO-60		412+-48.8 D/S/ML
818			33993		RU-106	A	
819			33993		SB-125	A	
820			33993		CS-134		43200+-1830 D/S/ML
821			33993		CS-137		744000+-43300 D/S/ML
822			7900		EPA-TOX		See Individual Elements Below
823			7906		SELENIUM		< 1.14893 mg/L
824			7900		EPA-TOX		See Individual Elements Below
			• •				

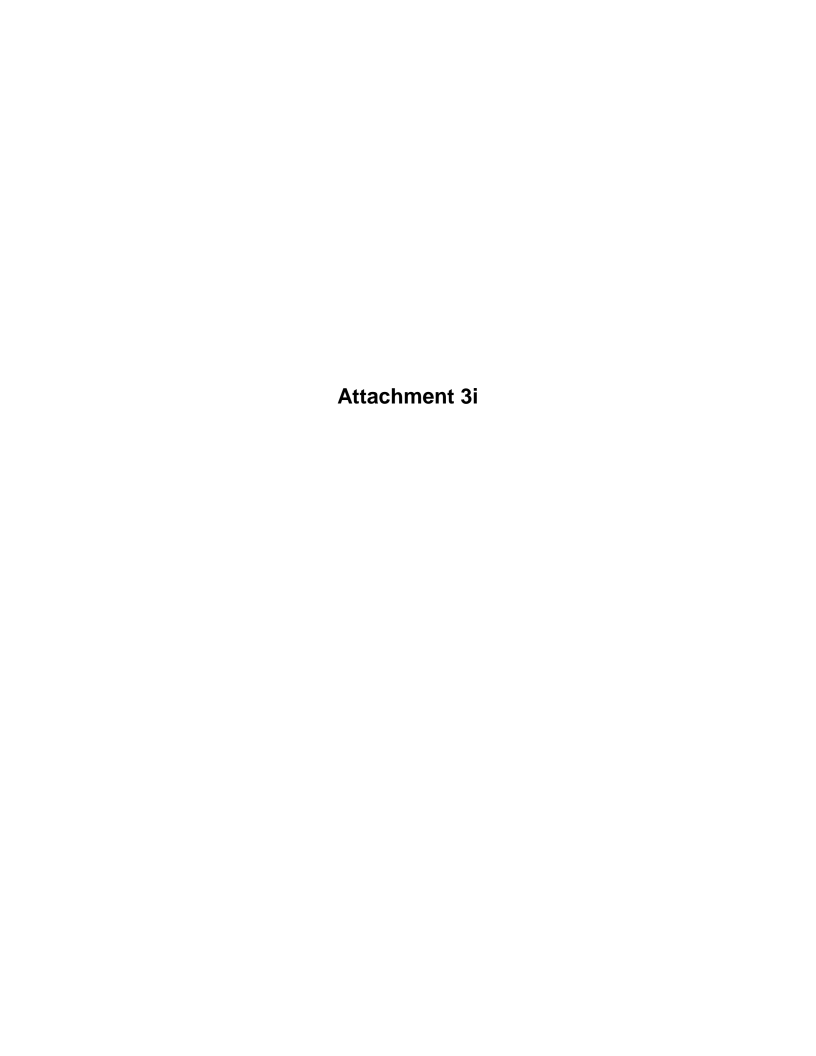
```
Log #
              Request Name
                               Log Type
                                             Charge Num Log Approval Info
  I
   d Lab
                        Meth a Ana-
   x ID
                              b lyst Analyte ARL
                         #
                                                               Result
 825 5
                        7906
                              7WDT SELENIUM C.. 3.70214+-.832765 mg/L
 843 60 CHK
                        7985
                              7CWL FLASHPOINT A.. NO Flash @ 60.0 deg C corrected
 844 END KNOWN
                        7985 7CWL FLASHPOINT C.. Flashed @ 60.0 deg C corrected
 855 PLASTIC 1
                      7100 7CBG METALS-ICP C.. See Individual Elements Below
                      7101 7CBG ALUMINUM A.. .618728+-3.79931E-02 MOLAR
 856 PLASTIC 1
                      7900 7VJJ EPA-TOX
 857 PLASTIC 1
                                               C.. See Individual Elements Below
 858 PLASTIC 1
                      7901 7VJJ ARSENIC
                                               C... < 6.93498 \text{ mg/L}
                    67093 7BKH FLUORIDE A.. 79.1488+-6.80393 UG/ML 7100 7CBG METALS-ICP C.. See Individual Elements Below
 860 PLASTIC 1
 861 PLASTIC 1
                      7105 7CBG IRON A.. 2.15423E-02+-2.18548E-03 MOLAR
                     7900 7WDT EPA-TOX
 862 PLASTIC 1
                                               C.. See Individual Elements Below
 863 PLASTIC 1
                      7906 7WDT SELENIUM C.. < 2.56205 mg/L
                   7900 7WDT EPA-TOX C.. See Individual Elements Below
7906 7WDT SELENIUM C.. < 2.56205 mg/L
77168 7MLE SULFATE A.. 2.00196E-03+-1.87812E-04 Molar
7900 7VJJ EPA-TOX C.. See Individual Elements Below
 864 PLASTIC 2
 865 PLASTIC 2
866 3
 915 PLASTIC 1R
                      7908 7SDN NICKEL
                                              C.. 6.98209+-2.56196 mg/L
 918 SPIKE
                      9304 9HCJ GCMS-2TRAP A.. SEE ATTACHED
 930505-17 WM-102:CH
                                                    13120-200-001940209 20:29 SPLMGT SPLMGT
                      7981 7 SP-GR
7981 7 SP-GR
67015 7MLE ACID
7920 7 URANIUM
 679 1
                                             C.. *(CANCELLED BY REQUESTOR)*
                                   SP-GR
                                  SP-GR
 680 2
                                               C.. *(CANCELLED BY REQUESTOR) *
 681 2
                                               A.. < .229527 NACID
                                             C.. *(CANCELLED BY REQUESTOR)*
 682 2
                                   URANIUM
 683 4
                       7972
                              7
                                   UDS
                                               C.. *(CANCELLED BY REQUESTOR) *
 684 2
                      57171
                              7MLE CHLORIDE A.. < 7.24282 UG/ML
                       7074 7
 685 3
                                               C.. *(CANCELLED BY REQUESTOR)*
                                   NITRATE
 686 2
                              7
                       7093
                                   FLUORIDE
                                               C.. *(CANCELLED BY REQUESTOR)*
                     7168 7 SULFATE C.. *(CANCELLED BY REQUESTOR)*
7961 7MLE RADIO CHEM C.. Prep Completed 050693 04:07
7961 7MLE SPEC CHEM C.. Prep Completed 050693 04:08
33993 3DAO GAMMA SCAN A.. ** No Nuclides Identified.
 687 3
 688 5
 689 PLASTIC 1
 690 5
                     23381 3BJS TOTAL SR A.. << 3.74185+-4.81351 D/S/ML
691 5
                      3998 3 TRANS-UA
692 5
                                               C.. *(CANCELLED BY REQUESTOR) *
                      3011 3
                                   TRITIUM C.. * (CANCELLED BY REQUESTOR) *
693 5
                    7100 7
                                   METALS-ICP C.. *(CANCELLED BY REQUESTOR)*
 694 PLASTIC 1
```

```
Log # Request Name Log Type Charge Num Log Approval Info
   Ι
   d Lab
                           Meth a Ana-
x ID
                                b lyst Analyte ARL
                            #
                                                                      Result
 930513-14 WM-102ET ????
                                                         35200-400-001951205 10:23 JEAN M CASTEEL
                           3281 3 NI-63 C. Entry Cancelled 3431 3 TECHNETIUM C. Entry Cancelled
 110 1
111 1
                          24900 4TEL URANIUM A.. * 6.54354E-03 G/KG
112 1
                    24900 4TEL URANIUM A. 6.54354E-U3 G/KG
24901 4TEL U-234 A. DATA NOT AVALIABLE
24902 4TEL U-235 A. DATA NOT AVALIABLE
24903 4TEL U-236 A. DATA NOT AVALIABLE
24904 4TEL U-238 A. DATA NOT AVALIABLE
24905 4TEL U FOR ACC. A. * 6.54354E-03 G/KG
17929 7BGP URAN. PREP C. 071993 09:26 TEL 071993 13:29
113 1
116 1
117 1
118 1
119 1
                       43204 3IDG NP-237 A.. < 137.824 D/S/ML
```

Log	#	Reques	t Name	Lo L	g Type	(	Charge Num I	Log Approval Info	
đ	Lab		Meth	a An	a-				
x	ID		#		st Analyte	ART.	Resu	1 <b>1</b> +	
							Repu		
120	1		3202	3IDG	PU-238	Α	3952.09+-326.	425 D/S/ML	_
121	1		3203	3IDG	PU-239	Α	< 688.675 D/S	S/ML	
122	1		3200	3	PU-240		Entry Cancelle		
123	1		3200	3	PU-241		Entry Cancelle		
124	1		3200	3	PU-242		Entry Cancelle		
125	1		3201	3IDG	AM-241		< 97.3643 D/S		
126	1		3201	3IDG	CM-244	Α	< 9.73E+1 D		
127	1		7961		RADIO CHEM			ed 062793 10:36	
475	4		2110		SODIUM	Α			
476	4		2190		POTASSIUM	Α		<del>-</del>	
477	4		7961		SP CHEM	С		ed 071693 23:04	
478	4		7100					Elements Below	
479	4		7101		ALUMINUM	c			
480			7102		BORON	Α			
481			7105		IRON	Α		884E-04 MOLAR	
482			7110		ZIRCONIUM	Α			
483			7111		CALCIUM	Α			
484			67093		FLUORIDE	Α			
485			7900		EPA-TOX			Elements Below	
486			7903		CADMIUM	A			
487			77168		SULFATE	Α		9.66199E-05 Mola	_
488			77074		NITRATE	Α		486E-02 Molar	<b>-</b>
489			57171		CHLORIDE		< 30.1508 UG/		
490			7972	7BGP			* 15 ÜG/ML	ML	
491			7980		SP-GR	A		017E-03 @ 25/4	
492			7980		SP-GR	A		017E-03 @ 25/4	
493			67015		ACID			•	
494			33993			A	** See Index	108E-02 NACID	**
495			13382	3 DAO	SR-89	C			^ ^
496			13382	3	SR-90	C	WRONG METHOD		
497			7961		SPEC CHEM	c	WRONG METHOD		
498			7100					d 071693 08:31 Elements Below	
499			7100						
500			67920		ALUMINUM	A		973E-02 MOLAR	
501					URANIUM	A		7.88716E-04 G/L	
502			33993		CS-134	A			
			33993		CS-137	A	577000+-23000		
503 504			23381		TOTAL SR	A	1.02623+149		
304	T		3993	3	GAMMA SCAN	С.,	ALREADY DONE		
9307	12-12	WM-102:					13120-200-0019	31128 02:58 SPLMO	т сртмат
243		2021	47981	<b>78ET</b>	SP-GR	Α			or Diffici
244	_		47981		SP-GR	A		495E-04 @ 25/4	
245			67015		ACID	A		967E-02 NACID	
246			7972	7KFM			NO VISIBLE S		
247			57171		CHLORIDE				
247			77074		NITRATE	A		523 UG/ML 568E-02 Molar	
249					FLUORIDE	A			
250			67093			A			_
Z30 .	J		77168	/KFM	SULFATE	Α	1.5/332E-U3+-	9.69884E-05 Molar	-

Log I	#	Reques	t Name	Log L	g Type	(	Charge Num Log Approval Info
đ	Lab		Meth	a Ana	<b>a~</b>		
x	ID		#	b lys	st Analyte	ARL	Result
251	5		7961	7KFM	RADIO CHEM	С	Prep Completed 071693 23:06
252	5		33993				** See Index 308 thru 310. **
253			33381	3KLJ	TOTAL SR	Α	<< 1.17428+-1.17347 UCI/ML
254	PLASTIC	1	7100	7SDN	METALS-ICP	С	See Individual Elements Below
255	PLASTIC	1	7101	7SDN	ALUMINUM	С	> .190546 MOLAR
256	PLASTIC	1	7102	7SDN	BORON	Α	< 3.75619E-03 MOLAR
257	PLASTIC	1	7105	7SDN	IRON	Α	3.36632E-02+-7.75552E-04 MOLAR
258	PLASTIC	1	7110	7SDN	ZIRCONIUM	Α	< 1.56506E-03 MOLAR
259	PLASTIC	1	7111	7SDN	CALCIUM	Α	< 2.54008E-03 MOLAR
	PLASTIC		7900	7SDN	EPA-TOX	С.,	See Individual Elements Below
	PLASTIC		7903	7SDN	CADMIUM	Α	28.4015+-1.8716 mg/L
	PLASTIC		7900	7SDN	EPA-TOX	С	See Individual Elements Below
263	PLASTIC	2	7903	7SDN	CADMIUM	Α	27.4401+-1.83477 mg/L
	PLASTIC		2110	2JSL	SODIUM	Α	=1.2E-02 Molarity
265	PLASTIC	1	2190	2JSL	POTASSIUM	Α	=9.3E-03 Molarity
	PLASTIC	_	7961	7KFM	SP CHEM M	С	Prep Completed 071693 23:06
305	PLASTIC	1	7100	7SDN	METALS-ICP	С	See Individual Elements Below
306	PLASTIC	1	7101	7SDN	ALUMINUM	Α	.912501+-3.90068E-02 MOLAR
307	1		7961	7CBG	SPEC CHEM	С	Prep Completed 071693 08:32
308	_		33993	3DAO	RU-106	Α	11900+-4280 D/S/ML
309	5		33993	3DAO	CS-134	Α	35800+-1750 D/S/ML
310			33993	3DAO	CS-137	Α	633000+-23400 D/S/ML
313	5		23381	3KLJ	TOTAL SR	Α	41826.1+-1479.44 D/S/ML

\*\*\*\*\*\*\*\* END \*\*\*\*\*\*



### **ENGINEERING DESIGN FILE**

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# <u>CPP-601 Deep Tanks – VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101</u>

Metals, Anions, and Miscellaneous

Metals, Anions, and Miscellaneous										
Analyte	Units	Method Number	Sample Le 981109		Sample Le 990119		Sample Log # 9902162		Sample Log # 9904133	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1	0.88		0.71	В	0.6	В	0.7	В
Acidity	N	AC7012								
Aluminum	μg/L	SW6010B	1.02E+05		6.09E+04	E	5.39E+04		1.81E+05	
Antimony	μg/L	SW6010B			1.04E+02	В	4.00E+01	ΒN	5.52E+01	В
Arsenic	µg/L	SW6010B	1.66E+01	U	2.52E+01	U	2.37E+01	U	2.37E+01	U
Barium	µg/L	SW6010B	1.09E+02		1.32E+02		1.34E+02		1.23E+02	
Beryllium	μg/L	SW6010B	< 1E00		2.3E00		2.3E00		2.7E00	
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	7.34E+02		1.48E+02		1.02E+02		3.85E+02	
Calcium	µg/L	SW6010B								
Chloride	μg/L	AC7171	1.06E+05				1.26E+05	В	8.37E+04	
Chromium	μg/L	SW6010B	1.89E+03		8.60E+02		1.94E+03		1.04E+03	
Cobalt	µg/L	SW6010B								
Copper	μg/L	SW6010B								
Fluoride	μg/L	AC7093	4.63E+04	В	2.55E+04	U	2.60E+04		2.69E+04	В
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	9.48E+02		8.15E+02		3.35E+02		3.79E+02	
Manganese	μg/L	SW6010B								
Mercury	μg/L	SW7470A	2.69E+03		1.44E+03		2.06E+03	N	5.62E+03	
Nickel	μg/L	SW6010B	8.52E+02		3.64E+02		8.17E+02		7.48E+02	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	2.00E+01		1.66E+01	U	2.67E+01	U	2.67E+01	U
Silver	μg/L	SW6010B	5.42E+01		6.4E00	U	2.14E+01	В	2.76E+01	В
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B			2.34E+01	U	2.84E+01	U	2.84E+01	U
Uranium	μg/L	AC7920	4.97E+02				3.09E+02		1.89E+02	U
Vanadium	μg/L	SW6010B			8.4E00	В	1.07E+01	В	1.42E+01	В
Zinc	μg/L	SW6010B			4.62E+03		2.20E+03		1.04E+03	
Zirconium	μg/L	SW6010B								
UDS	µg/L	AC7972	1.00E+05		3.00E+05		4.04E+04	В	4.57E+04	
TIC	μg/Ł	AC8060					4.66E+04	U	1.16E+04	U
TOC	µg/L	SW9060	2.36E+05	В	1.25E+05		8.76E+04	В	1.36E+05	

### **ENGINEERING DESIGN FILE**

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# CPP-601 Deep Tanks - VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)

Metals, Anions, and Miscellaneous (con't.)

Analyte	Units	Method Number	Sample Lo 990421		Sample Lo 990610		Sample Log # 9907081		Sample Log # 9908101	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1	0.7	U						
Acidity	N	AC7012			3.0E00	N	3.5E-01	В	2.87E-01	В
Aluminum	μg/L	SW6010B	5.08E+05		8.74E+04		6.06E+04		2.02E+05	
Antimony	μg/L	SW6010B	9.28E+01	В	4.90E+01	В	1.49E+01	В	6.34E+01	В
Arsenic	μg/L	SW6010B	2.37E+01	В	1.48E+01	U	2.18E+01	В	5.96E+01	В
Barium	μg/L	SW6010B	7.00E+01		9.92E+01		9.65E+01		7.04E+01	
Beryllium	μg/L	SW6010B	4.2E00		4.9E00		1.1E00		2.8E00	
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	2.96E+02		1.16E+02		1.86E+02		1.94E+02	
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171					,			
Chromium	µg/L	SW6010B	6.75E+02	N	1.28E+03		5.72E+02		5.71E+02	
Cobalt	μg/L	SW6010B	3.55E+01				4.18E+01		6.55E+01	
Copper	μg/L	SW6010B	4.62E+02				1.07E+03		8.04E+02	
Fluoride	µg/L	AC7093	2.56E+04	U	1.32E+04	В	1.87E+04	В	1.82E+04	В
Iron	μg/L	SW6010B								
Lead	µg/L	SW6010B	4.90E+02		3.03E+02		2.67E+02		4.42E+02	
Manganese	μg/L	SW6010B	1.53E+03				6.72E+02		1.10E+03	
Mercury	μg/L	SW7470A	6.97E+03		2.73E+03		1.22E+03		4.33E+03	Е
Nickel	μg/L	SW6010B	4.09E+02	N	5.39E+02	E	3.66E+02		4.72E+02	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	2.17E+01	U	2.17E+01	U	2.17E+01	U	2.17E+01	U
Silver	µg/L	SW6010B	3.23E+01	В	1.24E+02	N	2.29E+01	В	1.76E+01	В
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	2.24E+01	U	2.24E+01	U	2.24E+01	U	2.24E+01	U
Uranium	μg/ <b>L</b>	AC7920	1.89E+02	U					1.94E+02	
Vanadium	μg/L	SW6010B	6.6E00	В	1.33E+01	В	7.6E00	В	4.3E00	В
Zinc	μg/L	SW6010B	1.48E+03		1.62E+03		9.48E+02		1.67E+03	
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972							2.7E+03	UE
TIC	μg/L	AC8060	2.23E+04	U					1.80E+04	U
TOC	μg/L	SW9060	9.88E+04	В					1.50E+05	

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## <u>CPP-601 Deep Tanks – VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)</u>

Metals, Anions, and Miscellaneous (con't.)										
Analyte	Units	Method Number	Sample Lo 991101		Sample L 991122		Sample Log # 9911301		Sample Log # 9912131	
***************************************			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1								
Acidity	N	AC7012	5.1E-01	U	5.0E-01	В	3.0E-01	В	5.0E-01	В
Aluminum	μg/L	SW6010B	6.19E+04		2.21E+04		2.06E+05		2.34E+04	
Antimony	μg/L	SW6010B	6.34E+01		3.9E00	U	2.05E+01	ΒN	2.04E+01	ΒN
Arsenic	μg/L	SW6010B	5.34E+01		3.9E00	U	4.2E00	U	1.28E+01	В
Barium	µg/L	SW6010B	1.32E+02		4.10E+02		2.19E+02		9.01E+01	
Beryllium	µg/L	SW6010B	2.41E+01		4.3E00		3.1E00		9.0E00	В
Boron	µg/L	SW6010B								
Cadmium	μg/L	SW6010B	9.46E+01		6.56E+01		9.01E+01		4.01E+01	
Calcium	µg/L	SW6010B								
Chloride	µg/L	AC7171								
Chromium	μg/L	SW6010B	8.22E+02		9.62E+02		5.14E+02		5.55E+02	
Cobalt	μg/L	SW6010B	7.74E+01		5.19E+01		3.71E+01		3.53E+01	
Copper	μg/L	SW6010B	8.59E+02	N	1.12E+03		3.96E+02		5.34E+02	
Fluoride	μg/L	AC7093	1.98E+04	В	1.68E+04	В	2.17E+04	В	1.56E+04	В
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	2.76E+03		6.82E+02		1.99E+02		9.83E+02	
Manganese	μg/L	SW6010B	8.33E+02		3.76E+02		8.99E+02		3.75E+02	
Mercury	μg/L	SW7470A	1.34E+03		7.08E+02		1.70E+03		5.57E+02	
Nickel	µg/L	SW6010B	5.29E+02		5.90E+02		3.62E+02		3.54E+02	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	1.23E+01	В	3.7E00	U	3.2E00	U	7.0E00	В
Silver	µg/L	SW6010B	6.61E+01	N	4.11E+02		2.40E+01		1.61E+01	В
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	µg/L	SW6010B	2.39E+01	В	4.5E00	U	6.8E00	В	4.6E00	U
Uranium	μg/L	AC7920	4.05E+02		2.03E+02		1.70E+02		2.94E+02	
Vanadium	μg/L	SW6010B	2.34E+01	В	9.6E00	В	2.1E00	U	1.23E+01	В
Zinc	μg/L	SW6010B	1.59E+03	E	6.79E+02		9.42E+02		6.56E+02	
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972								
TIC	μg/L	AC8060	4.79E+03	В	3.5E+03	U	2.76E+04	U	1.38E+04	U
TOC	μg/L	SW9060	1.14E+05		4.34E+04		1.87E+05		6.64E+04	

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### <u>CPP-601 Deep Tanks – VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)</u>

Metals, Anions, and Miscellaneous (con't.)

Analyte	Units	Method Number	Sample Lo 000110		Sample Lo		Sample L 000214		Sample Log # 0003202	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1								
Acidity	N	AC7012	6.0E-01		3.0E-01	В	7.0E-01		4.0E-01	В
Aluminum	μg/L	SW6010B	2.28E+04		6.64E+04		4.83E+04		1.41E+05	
Antimony	μg/L	SW6010B	1.79E+01	В	2.58E+01	В	1.17E+01	ΒN	1.91E+01	ΒN
Arsenic	μg/L	SW6010B	1.28E+01	В	4.6E00	В	5.8E00	В	5.8E00	В
Barium	μg/L	SW6010B	2.23E+02	E	1.03E+02	E	2.65E+02		6.85E+02	
Beryllium	μg/L	SW6010B	7.8E00		3.4E00		9.1E00		2.8E00	
Boron	μg/L	SW6010B								
Cadmium	μg/L	SW6010B	4.66E+01		4.15E+01		4.20E+01		1.97E+02	
Calcium	μg/L	SW6010B								
Chloride	µg/L	AC7171								
Chromium	μg/L	SW6010B	7.03E+02	N	4.31E+02		8.75E+02		1.01E+03	
Cobalt	μg/L	SW6010B	4.28E+01		1.81E+02		8.89E+01		7.33E+01	
Copper	μg/L	SW6010B	6.88E+02	N	2.90E+02		4.31E+02		1.13E+03	
Fluoride	μg/L	AC7093	1.53E+04	В	1.52E+04		1.74E+04	В	2.85E+04	В
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	8.65E+02		1.34E+02		6.52E+02		6.30E+02	
Manganese	μg/L	SW6010B	3.71E+02		4.19E+02		5.68E+02		1.62E+03	
Mercury	μg/L	SW7470A	7.59E+02		9.29E+02		2.51E+02		3.07E+03	E
Nickel	μg/L	SW6010B	4.70E+02	N	2.97E+02		5.81E+02		7.83E+02	
Nitrate	μg/L	AC7074								
Phosphorus	µg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	4.5E00	В	3.2E00	U	3.7E00	U	3.7E00	U
Silver	μg/ <b>L</b>	SW6010B	4.2E00	В	7.2E00	В	4.01E+01		1.39E+01	
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	1.16E+01	В	1.10E+01	В	4.5E00	U	4.5E00	U
Uranium	μg/L	AC7920	2.87E+02		1.40E+02	U	1.88E+02		2.10E+02	
Vanadium	μg/L	SW6010B	1.07E+01		5.2E00	В	1.28E+01	В	5.5E00	В
Zinc	µg/L	SW6010B	6.32E+02		6.29E+02	Е	1.21E+03		9.42E+03	
Zirconium	μg/L	SW6010B								
UDS	µg/Ł	AC7972					5.75E+04		1.90E+05	
TIC	µg/L	AC8060	9.40E+03	U	2.81E+04	U	1.41E+04	U	2.81E+04	UE
тос	μg/L	SW9060	5.39E+04		8.15E+04		9.57E+04		2.65E+05	

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#### <u>CPP-601 Deep Tanks – VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)</u>

Volatile Organic Compounds and Semi-volatile Organic Compounds

Analyte	Units	Method Number	Sample L 981109		Sample L 990119		Sample L 99021		Sample Log # 9904133	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
Semi-Volatile O	rganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	3.2E+01	м
2,4-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	υ
4-Nitrophenol	µg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	U
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	U M
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	1.5E+02	D	2.5E+01	U	3.05E+02	D	7.0E+01	
Butylbenzyl phthalate	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	υ
Diethylphthalate	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	UM
Di-n-octyl phthalate	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	U	2.5E+01	U
Nitrobenzene	μg/ <b>L</b>	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	บ	2.5E+01	U M
n- Nitrosodimethylamine	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2.5E+01	υ	2.5E+01	U M
Pyridine	μg/L	SW8270C	3E+01		2.5E+01	U	2.5E+01	U	2.5E+01	U
Tri-n-butyl phosphate	μg/L	SW8270C	1.3E+04	D	7.1E+03	D	2.5E+01	U	2.5E+01	UM
Volatile Organic	Compo	unds								
1,1-Dichloroethane	µg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	U
2-Butanone	μg/L	SW8260A	4.0E00	U	4.0E00	U	4.2E00		1.0E+01	υ
2-Hexanone	µg/L	SW8260A	3.0E00	U	3.0E00	U	1.0E00	U	1.0E+01	UM
4-Methyl-2-pentanone	µg/L	SW8260A	7.8E00	J	4.0E00	U	1.0E00	U	1.0E+01	บ
Acetone	μg/L	SW8260A	1.6E+02		3.9E+01		1.0E+01	U	2.6E+02	E
Benzene	μg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	υм
Bromodichloromethane	μg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	U
Bromoform	µg/L	SW8260A	3.0E00	U	3.0E00	U	1.0E00	U	1.0E+01	U
Bromomethane	μg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	U
Carbon disulfide	μg/L	SW8260A	2.0E00	υ	2.0E00	U	1.0E00	U	1.0E+01	U
Carbon tetrachloride	μg/L	SW8260A	2.0E00	U	2.0E00	U	1.3E00	J	1.0E+01	U
Chloroform	µg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	U
Chloromethane	µg/L	SW8260A	3.0E00	U	3.0E00	U	2.0E00	U	1.0E+01	υ
Dibromochloromethane	µg/L	SW8260A	2.0E00	U	2.0E00	U	1.0E00	U	1.0E+01	U
Dichloromethane	µg/L	SW8260A	2.0E00	IJ	2.0E00	U	1.0E00	U	1.0E+01	UM
Styrene	µg/L	SW8260A	3.0E00	U	3.0E00	U	1.0E00	U	1.0E+01	UN
Trichlorofluoromethane	μg/L	SW8260A							1.0E+01	U

Tentatively Identified Organic Compounds

no. Ided / total conc.	µa/L	21 / 1.87E+03
no. Idea / total conc.	µg/L	 

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#### <u>CPP-601 Deep Tanks – VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)</u>

Volatile Organic Compounds and Semi-volatile Organic Compounds (con't.)

Analyte	Units	Method Number	Sample I 99042		Sample L 99061		Sample L 99070		Sample L 99081	
		· · · · · · · · · · · · · · · · · · ·	Results	LQF	Results	LQF	Results	LQF	Results	LQI
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	µg/L	SW8270C	2E+01	UM	2E+01	UM	2.5E+01	UM	2.5E+01	UN
2,4-Dinitrotoluene	μ <b>g/L</b>	SW8270C	2E+01	U	2E+01	U	2.5E+01	U	2.5E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	2E+01	U	2E+01	U	2.5E+01	U	2.5E+01	U
1-Nitrophenol	μg/L	SW8270C	2E+01	UM	2E+01	U	2.5E+01	U	2.5E+01	U
1,6-Dinitro-2- nethylphenol	µg/L	SW8270C	2E+01	υм	2E+01	U M	2.5E+01	U	2.5E+01	UN
3is-(2-ethylhexyl) ohthalate	µg/L	SW8270C	2E+01	U	6.8E+01		5E+02	D	3.0E+02	D
Butylbenzyl phthalate	μg/L	SW8270C	2E+01	U	2E+01	U	2.5E+01	U	2.5E+01	U
Diethylphthalate	μg/L	SW8270C	2E+01	U <b>M</b>	2E+01	U	2.5E+01	U	2.5E+01	UN
Di-n-octyl phthalate	μg/L	SW8270C	2E+01	U	2E+01	U	2.5E+01	U	2.5E+01	U
Nitrobenzene	μg/L	SW8270C	2E+01	UM	2E+01	UM	2.5E+01	UM	2.5E+01	U
n- Nitrosodimethylamine	μg/L	SW8270C	2E+01	U	2E+01	U	2.5E+01	UM	2.5E+01	υ
yridine 2	μg/L	SW8270C	2E+01	U	2E+01	U	2.5E+01	U	1.6E+03	D
ri-n-butyl phosphate	μg/L	SW8270C	1.5E+02	М	2E+01	U	9.6E+03	D	1.4E+04	DI
/olatile Organic	Compo	unds ·								
,1-Dichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	UM	1E+01	U
,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
2-Butanone	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
-Hexanone	μg/L	SW8260A	1E+01	UM	1E+01	UM	1E+01	UM	1E+01	U
-Methyl-2-pentanone	µg/L	SW8260A	1.2E+01		1E+01	U	1E+01	U	1E+01	U
Acetone	μg/L	SW8260A	1E+03	E	1E+01	UM	1E+01	UM	5.5E+02	E
Benzene	μg/L	SW8260A	1E+01	UM	1E+01	UM	1E+01	UM	1E+01	UN
Bromodichloromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Bromoform	μg/L	SW8260A	1E+01	U	1E+01	UM	1E+01	U	1E+01	U
Bromomethane	μg/L	SW8260A	1E+01	U	4E00	JBM	2E00	JBM	3E00	JB
Carbon disulfide	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	υ
Carbon tetrachloride	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	υ
Chloroform	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Chloromethane	μg/L	SW8260A	1E+01	U	1E+01	υz	1E+01	U	1E+01	U
Dibromochloromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Dichloromethane	μg/L	SW8260A	1E+01	U M	1E+01	U	1E+01	UM	1E+01	U
Styrene	μg/L	SW8260A	1E+01	U M	1E+01	U	1E+01	UM	1E+01	U
Trichlorofluoromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U

no. Ided / total conc. µg/L 9 / 3.28E+02 15 / 8.13E+02 22 / 1.47E+03 7 / 6.26E+02

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#### <u>CPP-601 Deep Tanks – VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)</u>

Volatile Organic Compounds and Semi-volatile Organic Compounds (con't.)

Analyte	Units	Method Number	Sample L 99110		Sample L 991122		Sample L 99113		Sample Log # 9912131	
			Results	LQF	Results	LQF	Results	LQF	Results	LQI
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	4E+01	UM	4E+01	UM	4E+01	UM	6E+01	UN
2,4-Dinitrotoluene	μg/ <b>L</b>	SW8270C	4E+01	U	2E+01	U	2E+01	U	2E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	4E+01	U	2E+01	U	2E+01	U	2E+01	U
4-Nitrophenol	μg/L	SW8270C	4E+01	υм	4E+01	U	4E+01	U	6E+01	U
4,6-Dinitro-2- nethylphenol	μg/L	SW8270C	4E+01	U	4E+01	UM	4E+01	U M	6E+01	U
3is-(2-ethylhexyl) ohthalate	μg/L	SW8270C	4.9E+01	М	1.2E+02		5E+01		2E+01	U
Butylbenzyl phthalate	μg/L	SW8270C	2.9E+01	JM	2E+01	U	2.4E+01		2E+01	U
Diethylphthalate	μg/L	SW8270C	4E+01	U	2E+01	UM	2E+01	υM	2E+01	UN
Di-n-octyl phthalate	μg/L	SW8270C	4E+01	UM	2E+01	υ	2E+01	U	2E+01	U
Nitrobenzene	µg/L	SW8270C	4E+01	U	2E+01	U	2E+01	U	2E+01	UM
n- Nitrosodimethylamine	μg/ <b>L</b>	SW8270C	4E+01	U	2E+01	U	2E+01	U	2E+01	UN
Pyridine	µg/L	SW8270C	4E+01	U	2E+01	U	1.3E+02		2E+01	U
ri-n-butyl phosphate	μg/L	SW8270C	8.8E+01	D	2E+01	UМ	2E+01	U M	6E+01	UN
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	1E+01	U	1.0E00	U	1.0E00	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	1.0E00	U	1.0E00	U	1E+01	U
2-Butanone	μg/L	SW8260A	1E+01	U	2.0E00	U	2.0E00	υ	1E+01	UN
2-Hexanone	µg/L	SW8260A	1E+01	U	2.0E00	U	2.0E00	U	1E+01	UN
1-Methyl-2-pentanone	μ <b>g/L</b>	SW8260A	1E+01	U	2.0E00	U	2.0E00	U	1E+01	UN
Acetone	μg/L	SW8260A	2.6E+02	E	1.3E+02	В	2.2E00	вјм	8.48E+02	ΕN
Benzene	μg/L	SW8260A	4E00	JBM	1.0E00	υM	1.0E00	U	1E+01	UN
Bromodichloromethane	μ <b>g/L</b>	SW8260A	1E+01	U	1.0E00	U	1.0E00	U	1E+01	U
Bromoform	μg/L	SW8260A	1E+01	U	2.0E00	U	2.0E00	U	1E+01	U
Bromomethane	μg/L	SW8260A	1E+01	U	2.0E00	U	2.0E00	U	1E+01	U
Carbon disulfide	μg/L	SW8260A	1E+01	U	1.0E00	U	1.0E00	U	1E+01	U
Carbon tetrachloride	µg/L	SW8260A	1E+01	U	2.0E00	U	2.0E00	U	1E+01	υ
Chloroform	µg/L	SW8260A	1E+01	U	1.0E00	U	1.0E00	UΖ	1E+01	U
Chloromethane	µg/L	SW8260A	1E+01	U	2.0E00	U	2.0E00	U	1E+01	UN
Dibromochloromethane	μg/L	SW8260A	1E+01	U	1.0E00	U	1.0E00	U	1E+01	U
Dichloromethane	μg/L	SW8260A	1E+01	UM	1.0E00	U M	1.0E00	UM	1E+01	UN
Styrene	μg/L	SW8260A	1E+01	UM	1.0E00	υм	1.0E00	UM	1E+01	U
		SW8260A	1E+01	U	1.0E00	U	1.0E00	U	1E+01	υ

no, Ided / total conc. μg/L 21/2.36E+03 27/1.37E+04 17/9.18E+02 8/1.20E+02

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#### CPP-601 Deep Tanks - VES-WG-100, VES-WG-101, VES-WH-100, & VES-WH-101 (con't.)

Volatile Organic Compounds and Semi-volatile Organic Compounds (con't.)

Analyte	Units	Method Number	Sample L 00011		Sample L 000111		Sample L 00021		Sample L 000320	
			Results	LQF	Results	LQF	Results	LQF	Results	LQ
Semi-Volatile Org	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	4E+01	UM	4E+01	UM	4E+01	U M	3.6E+01	JM
2,4-Dinitrotoluene	μg/L	SW8270C	4E+01	U	4E+01	U	4E+01	U	4E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	4E+01	U	4E+01	U	4E+01	U	4E+01	U
4-Nitrophenol	μg/L	SW8270C	4E+01	U	4E+01	U	4E+01	UM	4E+01	U I Z
4,6-Dinitro-2- methylphenol	μ <b>g/L</b>	SW8270C	4E+01	UM	4E+01	UM	4E+01	UM	4E+01	U I Z
Bis-(2-ethylhexyl) phthalate	μg/L	SW8270C	3.2E+01	J	5.6E+01		2.9E+01	J M	4.4E+01	м
Butylbenzyl phthalate	µg/L	SW8270C	7E00	J	1.5E+01	J	4E+01	U	4.1E+02	D
Diethylphthalate	μg/L	SW8270C	4E00	J	3E00	J	4E+01	U	8E00	JZ
Di-n-octyl phthalate	μg/L	SW8270C	4E+01	U	4E+01	U	4E+01	UM	4E+01	U N Z
Nitrobenzene	µg/L	SW8270C	4E+01	U	4E+01	U	4E+01	U	4E+01	U Z
n- Nitrosodimethylamine	μg/L	SW8270C	4E+01	U	4E+01	U	4E+01	U	4E+01	U:
Pyridine	μ <b>g/L</b>	SW8270C	4E+01	U	4E+01	U	4E+01	υ	7.2E+02	D
Fri-n-butyl phosphate	μg/L	SW8270C	8.5E+02	D	8.7E+02	D	1.4E+02		1.4E+03	ΕD
Volatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E+01	υ
2-Butanone	μg/L	SW8260A	2E00	UM	2E00	UM	1E+01	UM	1E+01	U
2-Hexanone	μg/L	SW8260A	2E00	UM	2E00	UM	1E+01	UM	1E+01	U
4-Methyl-2-pentanone	μg/L	SW8260A	2E00	UM	7E00	JM	1E+01	U	1E+01	U
Acetone	μg/L	SW8260A	1.6E+02	ВМ	2.5E+02	E M	1E+01	U B M	3.8E+02	E E M
Benzene	μ <b>g/L</b>	SW8260A	1E00	UM	1E00	UM	1E+01	UM	1E+01	UN
Bromodichloromethane	μ <b>g/L</b>	SW8260A	1E00	U	1E00	U	1E+01	U	1E+01	U
Bromoform	μg/L	SW8260A	2E00	U	2E00	U	1E+01	U	1E+01	U
Bromomethane	µg/L	SW8260A	2E00	U	2E00	U	1E+01	U	1E+01	U
Carbon disulfide	µg/L	SW8260A	1E00	υ	1E00	U	1E+01	U	1E+01	U
Carbon tetrachloride	μ <b>g/L</b>	SW8260A	2E00	U	2E00	U	1E+01	U	1E+01	u
Chloroform	µg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E+01	U
Chloromethane	µg/L	SW8260A	2E00	υM	2E00	UM	1E+01	U	1E+01	UI
Dibromochloromethane	µg/L	SW8260A	1E00	U	1E00	U	1E+01	U	1E+01	U
Dichloromethane	μg/L	SW8260A	1E00	UM	1E00	UM	2E00	JBM	1E+01	U
Styrene	μg/L	SW8260A	1E00	UM	1E00	UM	1E+01	UM	1E+01	UI
		SW8260A	1E00	υ	1E00	U	1E+01	U.	1E+01	u

14 / 5.88E+02

no. Ided / total conc. μg/L

22 / 6.54E+02

19 / 1.35E+03

21 / 8.09E+03



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#### PEWE Condensate Tanks - WL-106, WL-107, & WL-163

Metals, Anions, and Miscellaneous

Analyte	Units	Method Number	Sample L 990324		Sample Le 990327		Sample L 990428		Sample L 991115	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
pН		EPA150.1	1.43	В	1.43	В	0.75	В		
Acidity	N	AC7012							4.4E-01	
Aluminum	µg/L	SW6010B	2.41E+04		2.53E+04		3.67E+04		3.86E+04	
Antimony	µg/L	SW6010B	2.2E+01	U	2.2E+01	U	1.4E+01	U	4.2E00	U
Arsenic	μg/L	SW6010B	2.37E+01	U	2.37E+01	U	1.48E+01		4.2E00	U
Barium	μg/L	SW6010B	1.3E00	U	1.5E00	В	4.1E00	В	1.47E+01	
Beryllium	µg/L	SW6010B	2.0E-01	U	2.0E-01	U	1.0E-01	В	2.0E-01	В
Boron	µg/L	SW6010B								
Cadmium	µg/L	SW6010B	2.1E00	U	2.1E00	U	1.3E00	U	5.0E-01	U
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171								
Chromium	μg/L	SW6010B	2.62E+01		2.15E+01	В	3E00	U	5.70E+01	
Cobalt	μg/L	SW6010B	2.2E00	U	2.2E00	U	2.7E00	U		
Copper	μg/L	SW6010B	4.4E00	U	6.4E00	В	7.7E00	В	5.17E+01	
Fluoride	μg/L	AC7093	1.73E+03		1.24E+03	В	2.55E+04	U	7.37E+03	
Iron	µg/L	SW6010B								
Lead	μg/L	SW6010B	2.02E+01	U	2.02E+01	U	1.97E+01	U	4.3E00	U
Manganese	μg/L	SW6010B	5.1E00		5.5E00		4.3E00	В	7.6E00	
Mercury	µg/L	SW7470A	3.69E+01		2.98E+03	N	1.14E+03		5.25E+03	E
Nickel	μg/L	SW6010B	2.00E+01	В	1.76E+01	В	4.6E00	В	3.57E+01	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	2.67E+01	U	2.67E+01	U	2.17E+01	U	3.2E00	U
Silver	μg/L	SW6010B	6.2E00	U	6.2E00	U	3.5E00	U	1.7E00	U
Sodium	µg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	µg/L	SW6010B	2.84E+01	U	2.84E+01	U	2.24E+01	U	4.6E00	U
Uranium	μg/L	AC7920			1.88E+02	U	1.90E+02	Ų	1.38E+02	U
Vanadium	μg/L	SW6010B	3.6E00	U	3.6E00	U	4.3E00	U	2.1E00	U
Zinc	μg/L	SW6010B	1.01E+02		2.60E+02		5.12E+01		1.94E+02	N
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972								
TIC	µg/L	AC8060			2.33E+03	U	4.7E+03	U	3.45E+03	U
TOC	μg/L	SW9060			3.12E+04		5.22E+04		2.77E+04	

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#### PEWE Condensate Tanks - WL-106, WL-107, & WL-163 (con't.)

Metals, Anions, and Miscellaneous (con't.)

Analyte	Units	Method Number	Sample Lo 000307		Sample Lo 001114		Sample Lo 010221		Sample Le 010601	
			Results	LQF	Results	LQF	Results	LQF	Results	LQF
рH		EPA150.1								
Acidity	N	AC7012	2.2E-01		1.9E-01	В	6E-01		5.41E-01	
Aluminum	μg/L	SW6010B	2.55E+04		4.49E+04		2.41E+04		3.55E+04	
Antimony	μg/L	SW6010B	3.9E00	U	6.7E00	U	2.7E00	IJ	4.9E00	В
Arsenic	µg/L	SW6010B	3.9E00	U	4E00	U	4.5E00	U	2.9E00	U
Barium	µg/L	SW6010B	1.30E+01		1.3E00	В	2.8E00	В	2.79E+01	
Beryllium	μg/L	SW6010B	1.0E-01	В	3E-01	В	2E-01	В	3E-01	В
Boron	μg/L	SW6010B								
Cadmium	µg/L	SW6010B	7.0E-01	В	5E-01	U	3E-01	В	4E-01	U
Calcium	μg/L	SW6010B								
Chloride	μg/L	AC7171								
Chromium	μg/L	SW6010B	2.21E+01		1.25E+01		5.61E+01		4.37E+01	
Cobalt	μg/L	SW6010B			1.2E00	U	9E-01	В	1.1E00	В
Copper	μg/L	SW6010B	3.4E00	В	1.47E+01		1.96E+02		1.97E+01	
Fluoride	μg/L	AC7093	4.27E+03		5.7E+03		1.14E+05	UN	3.65E+04	U
Iron	μg/L	SW6010B								
Lead	μg/L	SW6010B	5E00	U	4.7E00	U	3.9E00	U	6.3E00	U
Manganese	μg/L	SW6010B	3.1E00		2.1E00		8E00		5.4E00	
Mercury	μg/L	SW7470A	6.52E+02		3.37E+04		4.41E+04		3.66E+04	
Nickel	μg/L	SW6010B	1.26E+01	В	8.2E00	В	3.36E+01		3.13E+01	
Nitrate	μg/L	AC7074								
Phosphorus	μg/L	SW6010B								
Potassium	μg/L	SW6010B								
Selenium	μg/L	SW6010B	3.7E00	U	4.8E00	U	2.9E00	U	4.8E00	U
Silver	μg/L	SW6010B	3.4E00	В	1.5E00	U	1.7E00	U	2E00	U
Sodium	μg/L	SW6010B								
Sulfur	μg/L	SW6010B								
Thallium	μg/L	SW6010B	4.5E00	U	4.2E00	U	3.8E00	U	4E00	U
Uranium	μg/L	AC7920	1.41E+02	U			3.2E+02	U	3.2E+02	U
Vanadium	µg/L	SW6010B	2.6E00	U	2.4E00	U	1.2E00	U	1E00	U
Zinc	μg/L	SW6010B	8.4E00		2.69E+01		2.26E+02		5.91E+01	
Zirconium	μg/L	SW6010B								
UDS	μg/L	AC7972	5.0E+03	U	5E+03	U	5E+03 TSS	U	1.07E+05 TDS	
TIC	μg/L	AC8060	1.41E+04	U	1.07E+04	UE	5.9E+03	UE	1.19E+04	UΕ
TOC	μg/L	SW9060	3.80E+04		5.92E+04	В	6.9E+04		5.43E+04	В

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#### PEWE Condensate Tanks - WL-106, WL-107, & WL-163 (con't.)

Volatile Organic Compounds and Semi-volatile Organic Compounds

Analyte	Units	Method Number	Sample L 990324		Sample L 990327		Sample L 99042		Sample Log # 9911151	
			Results	LQF	Results	LQF	Results	LQF	Results	LQI
Semi-Volatile Org	ganic C	ompounds								
2,4-Dinitrophenol	μg/L	SW8270C	3.6E+01	М	2.5E+01	UM	2.2E+01	М	6.6E+01	М
2,4-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
2,6-Dinitrotoluene	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
4-Nitrophenol	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	UM	4E+01	U
1,6-Dinitro-2- nethylphenol	μg/L	SW8270C	2.5E+01	UM	2.5E+01	UM	2E+01	UM	4E+01	UN
Bis-(2-ethylhexyl) ohthalate	µg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	3.1E+01	
Butylbenzyl phthalate	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
Diethylphthalate	μg/L	SW8270C	2.5E+01	UM	2.5E+01	UM	2E+01	UM	2E+01	UM
Di-n-octyl phthalate	μg/L	SW8270C	2.5E+01	UΖ	2.5E+01	υz	2E+01	U	2E+01	υ
Nitrobenzene	μg/L	SW8270C	2.5E+01	UM	2.5E+01	UM	2E+01	UM	2E+01	U
n- Nitrosodimethylamine	μ <b>g/L</b>	SW8270C	2.5E+01	UM	2.5E+01	Ų M	2E+01	U	2E+01	U
Pyridine	μg/L	SW8270C	2.5E+01	U	2.5E+01	U	2E+01	U	2E+01	U
ri-n-butyl phosphate	μ <b>g/L</b>	SW8270C	2.5E+01	UM	2.5E+01	UM	2E+03	М	2E+01	UN
/olatile Organic	Compo	unds								
1,1-Dichloroethane	μg/L	SW8260A	1E+01	U	1E+01	UΖ	1E+01	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	1E+01	UΖ	1E+01	U	1E+01	U
2-Butanone	µg/L	SW8260A	1.2E+01	М	1E+01	U M Z	8E00	J.	8E00	J
2-Hexanone	μg/L	SW8260A	1E+01	UM	1E+01	U M	1E+01	U	1E+01	U
4-Methyl-2-pentanone	μg/L	SW8260A	3E00	J	1E+01	U	1E+01	U	1E+01	U
Acetone	μg/L	SW8260A	2E+02	М	1.7E+02	ΜZ	2E+02	М	1.6E+02	
Benzene	μg/L	SW8260A	1E+01	U	1E+01	บ	1E+01	UM	6E00	JM
Bromodichloromethane	μg/L	SW8260A	1.1E+01		1E+01	υ	1E+01	U	1E+01	U
Bromoform	μg/L	SW8260A	2E00	J	1E+01	U	1E+01	U	1E+01	U
Bromomethane	μg/L	SW8260A	1E+01	U	3E00	JΖ	2.7E+01		1.4E+01	
Carbon disulfide	μg/L	SW8260A	2E00	J	1E+01	UΖ	1E+01	U	1E+01	U
Carbon tetrachloride	μg/L	SW8260A	5E00	J	1E+01	υ	1E+01	U	1E+01	U
Chloroform	μg/L	SW8260A	2.3E+01		1E+01	UΖ	1E+01	U	1E+01	U
Chloromethane	µg/L	SW8260A	1E+01	U	8E00	JΖ	7.0E+01		1E+01	U
Dibromochloromethane	µg/L	SW8260A	7E00	J	1E+01	U	1E+01	U	1E+01	U
Dichloromethane	μ <b>g/L</b>	SW8260A	1E+01	U	3E00	JΖ	1E+01	บ	1E+01	U
Styrene	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
							1E+01	U		UN

no. Ided / total conc. µg/L 13 / 1.29E+03 18 / 1.56E+03 19 / 5.46E+02 22 / 1.23E+03

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#### PEWE Condensate Tanks - WL-106, WL-107, & WL-163 (con't.)

Volatile Organic Compounds and Semi-volatile Organic Compounds (con't.)

Analyte	Units	Method Number	Sample Lo 000307		Sample Lo 001114		Sample L 010221		Sample Log # 0106011	
			Results	LQF	Results	LQF	Results	LQF	Results	LQI
Semi-Volatile Or	ganic C	ompounds								
2,4-Dinitrophenol	µg/L	SW8270C	1.3E+01	J	3.3E+01	М	4E+01	D D	2.8E+01	М
2,4-Dinitrotoluene	µg/L	SW8270C	2E+01	U	2E+01	υ	4E+01	ΟŲ	2E+01	U
2,6-Dinitrotoluene	μ <b>g/L</b>	SW8270C	2E+01	U	2E+01	U	4E+01	αυ	2E+01	U
4-Nitrophenol	μg/L	SW8270C	2E+01	U	2E+01	U	4E+01	UD	2E+01	U
4,6-Dinitro-2- methylphenol	μg/L	SW8270C	5E00	J	2E+01	U	4E+01	U <b>M</b> D	2E+01	UN
Bis-(2-ethylhexyl) ohthalate	μg/L	SW8270C	3.1E+01		6E00	J	2.2E+01	JÐ	2.4E+01	
Butylbenzyl phthalate	μg/L	SW8270C	2E+01	U	2E+01	U	4E+01	UD	2E+01	U
Diethylphthalate	μg/L	SW8270C	2E+01	UM	2E+01	UМ	4E+01	U M Z D	2E+01	UN
Di-n-octyl phthalate	µg/L	SW8270C	2E+01	U	2E+01	U	4E+01	U D	2E+01	U
Nitrobenzene	µg/L	SW8270C	2E+01	UM	2É+01	UM	4E+01	UD	2E+01	UN
n- Nitrosodimethylamine	μg/L	SW8270C	2E+01	UM	1.5E+02		2E+02	D	4.5E+02	Q
Pyridine	μg/L	SW8270C	2E+01	U	2E+01	U	4E+01	UD	2E+01	U
Tri-n-butyl phosphate	μg/L	SW8270C	1.7E+02	ΕM	6.6E+02	D M	4.4E+02	DE	3.4E+02	DN
Volatile Organic										
1,1-Dichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
1,1,1-Trichloroethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
2-Butanone	µg/L	SW8260A	5E00	J M	1E+01	U	1E+01	U	5E00	J
2-Hexanone	μg/L	SW8260A	1E+01	UM	1E+01	U	1E+01	υ	1E+01	U
4-Methyl-2-pentanone	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Acetone	μg/L	SW8260A	1.9E+02	вм	7.2E+01		5.3E+01		8.5E+01	
Benzene	μg/L	SW8260A	1E+01	UM	1E+01	U	1E+01	U	1E+01	U
Bromodichloromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Bromoform	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	υ	1E+01	U
Bromomethane	µg/L	SW8260A	1E+01	U	7E00	J	3.8E+01		9E00	J
Carbon disulfide	μg/L	SW8260A	1E+01	U	1E+01	υ	1E+01	U	1E+01	U
Carbon tetrachloride	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Chloroform	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	υ	1E+01	U
Chloromethane	μg/L	SW8260A	3E00	J	1E+01	UM	5.6E+01	М	2.3E+01	М
Dibromochloromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Dichloromethane	μg/L	SW8260A	1E+01	UM	1E+01	U	1E+01	U	1E+01	U
Styrene	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Trichlorofluoromethane	μg/L	SW8260A	1E+01	U	1E+01	U	1E+01	U	1E+01	U
Tentatively Ident									,	
		J	24 / 7.18E+03				13 / 3.64E+02		20 / 4.74E+02	SVO



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#### LET&D Bottoms Tank - WLL-195

Metals, Anions, and Miscellaneous

Analyte	Units	Method Number	Sample L 990327		Sample Le 990504		Sample Log # 0003285		
			Results	LQF	Results	LQF	Results	LQF	
pН		EPA150.1	0.34	U					
Acidity	N	AC7012			1.21E+01		1.20E+01		
Aluminum	μg/L	SW6010B	1.11E+06	5.11	1.72E+06		1.71E+06		
Antimony	μg/L	SW6010B	4.77E+01	B N E	1.66E+02	В	3.9E+01	U	
Arsenic	μg/L	SW6010B	3.57E+01	В	2.89E+02	В	3.9E+01	U	
Barium	μg/L	SW6010B	2.34E+01		2.45E+02		4.30E+01	В	
Beryllium	μg/L	SW6010B	4.6E00		6.6E00	В	1.10E+01		
Boron	μg/L	SW6010B							
Cadmium	μg/L	SW6010B	1.58E+01	В	3.30E+01	В	1.50E+01	В	
Calcium	μg/L	SW6010B							
Chloride	μg/L	AC7171							
Chromium	μg/L	SW6010B	3.54E+04		1.77E+05		5.28E+04		
Cobalt	μg/L	SW6010B	3.58E+03		2.13E+04		6.04E+03		
Copper	μg/L	SW6010B	1.86E+03		1.15E+04		2.92E+03		
Fluoride	μg/L	AC7093	9.93E+04	В	6.58E+04	В	2.43E+05	U	
Iron	μg/L	SW6010B							
Lead	μg/L	SW6010B	2.02E+01	U	2.17E+02	U	5.0E+01	U	
Manganese	μg/L	SW6010B	1.66E+03		6.08E+03		1.87E+03		
Mercury	μg/L	SW7470A	3.83E+04		5.55E+04		3.64E+04	BE	
Nickel	µg/L	SW6010B	4.42E+04		2.27E+05		7.0E+04		
Nitrate	μg/L	AC7074							
Phosphorus	μg/L	SW6010B							
Potassium	μg/L	SW6010B							
Selenium	μg/L	SW6010B	2.67E+01	U	2.39E+02	U	3.7E+01	U	
Silver	μg/L	SW6010B	6.2E00	UE	3.85E+01	UN	1.70E+01	В	
Sodium	μg/L	SW6010B							
Sulfur	μg/L	SW6010B							
Thallium	μg/L	SW6010B	2.84E+01	U	2.79E+02	В	4.5E+01	U	
Uranium	µg/L	AC7920	1.89E+02	U	1.89E+02	U	2.77E+01		
Vanadium	µg/L	SW6010B	8.49E+01		2.45E+02	В	2.6E+01	U	
Zinc	μg/L	SW6010B	8.80E+02	Ε	1.37E+03	E	6.96E+02	E	
Zirconium	μg/L	SW6010B							
UDS	µg/L	AC7972					2.22E+04		
TIC	μg/L	AC8060	2.46E+03		8.70E+04		1.41E+02	U	
TOC	μg/L	SW9060	1.41E+04		4.35E+03	В	7.32E+04	В	

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#### LET&D Bottoms Tank - WLL-195 (con't.)

Volatile Organic Compounds and Semi-volatile Organic Compounds

Analyte	Units	Method Number	Sample L 99032		Sample L 99050		Sample L 000328	
			Results	LQF	Results	LQF	Results	LQF
emi-Volatile Or	ganic C	ompounds						
,4-Dinitrophenol	μg/L	SW8270C	2.5E+03	U M D	2E+01	UM	1E+03	U M
,4-Dinitrotoluene	μg/L	SW8270C	2.5E+03	U D	2E+01	U	1E+03	U
6-Dinitrotoluene	μg/L	SW8270C	2.5E+03	U D	2E+01	U	1E+03	U
l-Nitrophenol I,6-Dinitro-2-	µg/L	SW8270C	2.5E+03	UD U M	2E+01	UM	1E+03	UM
ethylphenol is-(2-ethylhexyl)	µg/L	SW8270C	2.5E+03	D	2E+01	UM	1E+03	UM
hthalate	μg/L	SW8270C	2.5E+03	QU	2E+01	U	1E+03	U
lutylbenzyl phthalate	μg/L	SW8270C	2.5E+03	QU	2E+01	U	1E+03	U
ethylphthalate	μg/L	SW8270C	2.5E+03	U M D	2E+01	UM	1E+03	U
i-n-octyl phthalate	μg/L	SW8270C	2.5E+03	UD	2E+01	U	1E+03	U
itrobenzene	μg/L	SW8270C	2.5E+03	U M D	2E+01	UM	1E+03	U
itrosodimethylamine	µg/L	SW8270C	2.5E+03	U M D	2E+01	U	1E+03	U
								U
yridine	μg/L	SW8270C	2.5E+03	UD U M	2E+01	U	1E+03	UM
ri-n-butyl phosphate	μg/L	SW8270C	2.5E+03	D	2.9E+01	M	1E+03	D
olatile Organic	Compo	unds						
1-Dichloroethane	μg/L	SW8260A			1E+01	UY	1.5E+02	U
1,1-Trichloroethane	μg/L	SW8260A			1E+01	UY	1.5E+02	U
-Butanone	μg/L	SW8260A			1E+01	UY	1.5E+02	UM
Hexanone	μg/L	SW8260A			1E+01	UY	1.5E+02	UM
Methyl-2-pentanone	μg/L	SW8260A			1E+01	U	1.5E+02	UM
etone	μg/L	SW8260A			1E+01	U M Y	9.8E+02	вм
enzene	μg/L	SW8260A			1E+01	UM	1.5E+02	U M
					1E+01	U	1.5E+02	U U
omodichloromethane	μg/L 	SW8260A						
omoform	µg/L	SW8260A			1E+01	UY	1.5E+02	U
omomethane	μg/L	SW8260A			6.3E+02	EY	4.2E+01	JM
arbon disulfide	µg/L	SW8260A			1E+01	UY	1.5E+02	U
arbon tetrachloride	µg/L	SW8260A			1E+01	UY	1.5E+02	U
hloroform	µg/L	SW8260A			1E+01	UY	1.5E+02	U
nloromethane	μg/L	SW8260A			1E+01	UY	5.9E+02	
bromochloromethane	μg/L	SW8260A			1E+01	UY	1.5E+02	U
chloromethane	μ <b>g/L</b>	SW8260A			1E+01	UY	1.5E+02	UM
yrene	μg/L	SW8260A			1E+01	UY	1.5E+02	UMZ
•							1E+01	U
richlorofluoromethane	µg/L	SW8260A	a. mala		6E00	JY	15701	U
entatively Ident	iiied Or	ganic Comp	ounas	0)/0.0				
ded / total conc.	μg/L		12 / 6.93E+05	SVOC only	28 / 1.36E+04		8 / 1.58E+04	



#### TEMPLATE REPORT for WG:

Log Type: \*\* PLANT \*\*

Report for : FUEL Mailstop : 5201

Mailstop : 5201 Phone Number : 3244

Date Received : Date completed : Time Received : Time completed :

GWA charged : 100333S01 Reviewed by

MSA mR/hr : 1 Signature \_\_\_\_\_

Hazard Index : 1E6 Lab QC/QA reviewed by

Quality Level: NO Signature \_\_\_\_\_

COMMENTS: SEND COPY OF FINAL REPORT TO DAVE OLSEN, MS 5111

Analysis	Sample	Method Analyst	Results
AL/F RATIO	WG	1023	
Acid	WG	7012	
Aluminum	WG	7100	
COMPOSITES	WG	7963	
00111 02 - 1 - 2	WG	7963	
Chloride	WG	7171	
Flash Point	WG	7985	
Fluoride	WG	7093	
GROSS BETA	WG	7970	
Mercury	WG	7802	
Nitrate	WG	7074	
Potassium	WG	2118	
Sodium	WG	2118	
Sulfate	WG	7168	
TOC	WG	8060	
TOC(r)	WG	8060	
U234	WG	8920	
U235	WG	8920	
U236	WG	8920	
U238	WG	8920	
URANIUM FOR ACC	WG	8920	
URANIUM.	WG	8920	
Uranium	WG	7920	
	WG	7920	
Uranium PreP	WG	7929	
рН	WG	7016	
End of Report	26 Analyse	s.	

#### TEMPLATE REPORT for WH:

Log Type: \*\* PLANT \*\*

Report for : FUEL Mailstop : 5201

Mailstop : 5201 Phone Number : 3244

Date Received: Date completed: Time Received: Time completed:

GWA charged : 100332524 Reviewed by

MSA mR/hr : 1 Signature \_\_\_\_\_

Hazard Index : 1E6 Lab QC/QA reviewed by

Quality Level: NO Signature \_\_\_\_

COMMENTS: SEND COPY OF FINAL REPORT TO DAVE OLSEN, MS 5111

Analysis	Sample	Method Analyst	Results
AL/F RATIO	WH	1023	
Acid	WH	7012	
Aluminum	WH	7100	
COMPOSITES	WH	7963	
	WH	7963	
Chloride	WH	7171	
Flash Point	WH	7985	
Fluoride	WH	7093	
GROSS BETA	WH	7970	
Mercury	WH	7802	
Nitrate	WH	7074	
Potassium	WH	2118	
Sodium	WH	2118	
Sulfate	WH	7168	
TOC	WH	8060	
TOC(r)	WH	8060	
U234	WH	8920	
U235	WH	8920	
U236	WH	8920	
U238	WH	8920	
URANIUM FOR ACC	WH	8920	
URANIUM.	WH	8920	
Uranium	WH	7920	
	WH	7920	
Uranium PreP	WH	7929	
рН	WH	7016	
End of Report	26 Analyse	s.	

## FINAL REPORT for WG:101 10/28/02 Log Type: \*\* PLANT \*\*

Report for : NWCF Mailstop : 5218 Log Number : 02-10283 Phone Number : 6-5456

Date Approved : Jan 16 2003 Time Approved : 13:46 Date Received : Oct 29 2002 Time Received : 13:20

Reviewed by KIMBERLY WHITEHEAD GWA charged : 100323H55

MSA mR/hr : 1 Signature \_\_\_

Hazard Index : >1E4 Laboratory QA Review

Signature \_\_\_\_ PCBs >50 ppm : NO

COMMENTS: SEND COPY OF FINAL REPORT TO DAVE OLSEN, MS 5111

	Lab Field	
Analysis	Spl ID Spl ID	Method Analyst Results
AL/F RATIO		11023 RAH Ratio Not Performed
Acid	2BQ49 WG:101	57012 RDW 3.93E-01 +- 8.4E-02 Normal Acid
Aluminum	2BQ49 WG:101	52900 DSL 2.44934E-03 Molar
Chloride	2BQ49 WG:101	57171  RDW  5.4E+01 +- 1.3E+01  ug/mL
Flash Point	2BQ49 WG:101	17985 AWO NO FLASH @ 60.00 deg C corrected
Fluoride	2BQ49 WG:101	57093 AWO Not Detected: MDL=26 ug/mL
GROSS BETA	2BQ49 WG:101	87970 RAH 8.98E+05 +- 5.2E+04 B/Min/ml
Mercury	2BQ49 WG:101	77802 RAH 1.65E+00 +- 2.2E-01 ug/ml
Nitrate	2BQ49 WG:101	97074 BGP 5.22E-03 +- 5.8E-04 Molar
Potassium	2BQ49 WG:101	52900 DSL 1.19173E-03 Molar
Sodium	2BQ49 WG:101	52900 DSL 1.08865E-02 Molar
Sulfate	2BQ49 WG:101	97168  BGP  5.8E+01 +- 1.8E+01  ug/ml
TOC	2BQ49 WG:101	18060 BGP Not Detected: MDL=9e+01 ug/ml
U234	2BQ49 WG:101	28920 DDJ 3.4E-03 WEIGHT %
U235	2BQ49 WG:101	28920 DDJ 4.919E-01 WEIGHT %
U236	2BQ49 WG:101	28920 DDJ 2.39E-02 WEIGHT %
U238	2BQ49 WG:101	28920 DDJ 9.94808E+01 WEIGHT %
URANIUM.	2BQ49 WG:101	28920 DDJ
Uranium	2BQ49 WG:101	17920 BGP 2.21E-03 +- 5.9E-04 G/L
	2BQ49 WG:101	17920 BGP 2.43E-03 +- 6.2E-04 G/L
Uranium PreP	2BQ49 WG:101	17929 AWO 1.0E+00 ml
End of Report	21 results.	

## FINAL REPORT for WG:101 8 Log Type: \*\* PLANT \*\*

Log Number : 02-04242 Phone Number : 3244 Report for : FUEL Mailstop : 5201

Date Approved : May 28 2002 Time Approved : 15:34 Date Received : Apr 25 2002

Time Received: 10:40

Reviewed by KIMBERLY WHITEHEAD GWA charged : 51R101327

MSA mR/hr : 1 Signature \_\_\_\_

Hazard Index : 1E6 Laboratory QA Review

PCBs >50 ppm : NO Signature \_\_\_\_

COMMENTS: SEND COPY OF FINAL REPORT TO DAVE OLSEN, MS 5111 - Use GWA

51R101327

	Lab Field	
Analysis	Spl ID Spl ID	Method Analyst Results
AL/F RATIO	2AO12 WG	11023 GDD Ratio Not Performed
Acid	2AO12 WG	57012 RAH 4.98E-01 +- 7.9E-02 Normal Acid
Aluminum	2AO12 WG	87100 GDD 4.37E-03 +- 3.0E-04 MOLAR
Chloride	2AO12 WG	57171 RAH 8.0E+01 +- 1.2E+01 ug/mL
Flash Point	2AO12 WG	17985 GDD NO FLASH @ 60.00 deg C corrected
Fluoride	2AO12 WG	57093 AWO Not Detected: MDL=45.37 ug/mL
GROSS BETA	2AO12 WG	87970 RAH 9.84E+04 +- 4.2E+03 B/Min/ml
Mercury	2A012 WG	87802 RAH 1.57E+00 +- 4.5E-01 ug/ml
Nitrate	2A012 WG	97074 GDD 2.887E+04 +- 7.6E+02 ug/mL
Potassium	2AO12 WG	12800 SDN 4.94E+01 ug/mL
Sodium	2AO12 WG	12800 SDN 2.55E+02 ug/mL
Sulfate	2AO12 WG	97168 BGP 6.9E+01 +- 1.8E+01 ug/ml
TOC	2AO12 WG	18060  BGP  9.5E+01 +- 2.8E+01  ug/ml
U234	2AO12 WG	28920 DDJ 2.5E+00 WEIGHT %
U235	2AO12 WG	28920 DDJ 8.0E+00 WEIGHT %
U236	2A012 WG	28920 DDJ 0.0E+00 WEIGHT %
U238	2AO12 WG	28920 DDJ 8.95E+01 WEIGHT %
URANIUM FOR ACC	2AO12 WG	28920 DDJ 2.9E-04 g/kg
URANIUM.	2AO12 WG	28920 DDJ 2.9E-04 g/kg
Uranium	2AO12 WG	17920  RAH < 3.94794E-04  G/L
	2AO12 WG	17920 RAH < 3.94794E-04 G/L
Uranium PreP	2A012 WG	17929 BGP 1.0E+00 ml
End of Report	22 results.	

# F I N A L R E P O R T for WH:101 Log Type: \*\* PLANT \*\*

Log Number : 02-03272 Phone Number : 3244 Report for : FUEL Mailstop : 5201

Date Approved : Jun 26 2002 Time Approved : 14:24 Date Received : Mar 28 2002

Time Received: 09:21

GWA charged : 51R101327 Reviewed by JEFF LAUG

MSA mR/hr : 10 Signature \_\_\_\_

Laboratory QA Review Hazard Index : 1E6

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS: SEND COPY OF FINAL REPORT TO DAVE OLSEN, MS 5111.

	Lab Field	
Analysis	Spl ID Spl ID	Method Analyst Results
AL/F RATIO	2AK33	11023 AWO Ratio Not Performed
Acid	2AK33	57012  AWO  3.24E-01 +- 7.3E-02  Normal Acid
Aluminum	2AK33	52900 LAM 2.12167E-03 Molar
Chloride	2AK33	57171  AWO  7.5E+01 +- 1.2E+01  ug/mL
Flash Point	2AK33	17985 GDD NO FLASH @ 60.00 deg C corrected
Fluoride	2AK33	57093 AWO Not Detected: MDL=23.53 ug/mL
GROSS BETA	2AK33	87970 BGP $8.07E+03 +- 4.4E+02$ B/Min/ml
Mercury	2AK33	77802 RAH 1.003E+00 +- 7.4E-02 ug/ml
Nitrate	2AK33	97074 GDD $1.710E+04 +- 3.5E+02 \text{ ug/mL}$
Potassium	2AK33	12800 SDN 55.2 ug/mL
Sodium	2AK33	12800 SDN 236. ug/mL
Sulfate	2AK33	97168  GDD  7.1E+01 +- 1.8E+01  ug/ml
TOC	2AK33	18060  BGP  1.30E+02 +- 2.8E+01  ug/ml
U234	2AK33	28920 DDJ 0.0E+00 WEIGHT %
U235	2AK33	28920 DDJ 4.011E+00 WEIGHT %
U236	2AK33	28920 DDJ 0.0E+00 WEIGHT %
U238	2AK33	28920 DDJ 9.599E+01 WEIGHT %
URANIUM FOR ACC	2AK33	28920 DDJ 3.29E-04 g/kg
URANIUM.	2AK33	28920 DDJ 3.29E-04 g/kg
Uranium	2AK33	17920  BGP < 3.94794E-04  G/L
	2AK33	17920  BGP < 3.94794E-04  G/L
Uranium PreP	2AK33	17929 BGP 1.0E+00 ml
End of Report	22 results.	

### FINAL REPORT for WH:101 N/A Log Type: \*\* PLANT \*\*

Report for : FUEL Mailstop : 5201 Log Number : 01-08297 Phone Number : 3244

Date Approved : Oct 10 2001 Time Approved : 10:16 Date Received : Aug 30 2001

Time Received: 12:37

GWA charged : 51F1052B3 Reviewed by JEFF LAUG

MSA mR/hr : 3.0 Signature \_\_\_\_

Laboratory QA Review Hazard Index : 1E5

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS: SEND COPY OF FINAL REPORT TO DAVE OLSEN, MS 5111 - Use GWA

51f10a20

Analysis	Lab Field Spl ID Spl ID	Method Analyst Results
AL/F RATIO	1CN57	11023 BCS Ratio Not Performed
Acid	1CN57	57012 RNR 3.62E-01 +- 1.1E-02 Normal Acid
Aluminum	1CN57	87100 BCS 1.07E-03 +- 2.3E-04 MOLAR
Chloride	1CN57	57171 RNR 6.18E+01 +- 5.1E+00 ug/mL
Flash Point	1CN52 WH-101	17985 BCS NO FLASH @ 60.00 deg C corrected
Fluoride	1CN57	57093 RAH Not Detected: MDL=69.88 ug/mL
GROSS BETA	1CN57	87970 BCS 1.688E+05 +- 9.1E+03 B/Min/ml
Mercury	1CN57	87802 RAH 7.6E-01 +- 1.2E-01 ug/ml
Nitrate	1CN52 WH-101	97074 BCS 3.412E-01 +- 8.0E-03 Molar
Potassium	1CN57	12800 SDN 43.8 ug/mL
Sodium	1CN57	12800 SDN 186. ug/mL
Sulfate	1CN57	97168  BCS  5.93E+01 +- 4.3E+00  ug/ml
TOC	1CN57	18060 BGP 1.290E+02 +- 9.0E+00 ug/ml
U234	1CN57	24900 DDJ 1.6E-01 WEIGHT %
U235	1CN57	24900 DDJ 6.37E+00 WEIGHT %
U236	1CN57	24900 DDJ 3.8E-01 WEIGHT %
U238	1CN57	24900 DDJ 9.31E+01 WEIGHT %
URANIUM FOR ACC	1CN57	24900 DDJ 1.28E-04 g/kg
URANIUM.	1CN57	24900 DDJ 1.28E-04 g/kg
Uranium	1CN57	17920 BCS < 3.24219E-04 G/L
	1CN57	17920 BCS < 3.24219E-04 G/L
Uranium PreP	1CN57	17929 BGP 1.0E+00 ml
End of Report		



#### FINAL REPORT for WL-103 T/D

Log Type: \*\* PLANT \*\*

Log Number : 96-100815 Phone Number : 6-3226 Report for : NEILL Mailstop : 5116

Date Approved : Oct 12 1996 Time Approved : 16:48 Date Received : Oct 08 1996

Time Received: 22:06

GWA charged : 522020702 Reviewed by CLAYNE GRIGG

Signature \_\_\_\_\_ MSA mR/hr : <0.1

Hazard Index : <1E4 Laboratory QA Review

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS:

	Lab Field	
Analysis	spl ID spl ID	Method Analyst Results
Acid	6DC01 WL-103	17012 RAH titrated less than .5ml
Chloride	6DC01 WL-103	57171 RAH < 1.25947E+01 ug/ml
Fluoride	6DC01 WL-103	87092 RAH 2.40E+01 +- 1.7E+00 ug/ml
SpGr	6DC01 WL-103	47981 BCS 9.98685E-01 +- 2.6E-04 @ 25/4
Sulfate	6DC01 WL-103	97168 BCS < 9.03652E+00 uG/mL
Uranium	6DC01 WL-103	17920 RAH 1.99E-02 +- 1.2E-03 G/L
	6DC01 WL-103	17920 RAH 2.02E-02 +- 1.2E-03 G/L
рН	6DC01 WL-103	87017 RLC 2.46E+00 +- 1.6E-01 pH
End of Report	8 results.	

#### FINAL REPORT for WL-104 T/D

Log Type: \*\* PLANT \*\*

Log Number : 96-100816 Phone Number : 6-3226 Report for : NEILL Mailstop : 5116

Date Approved : Dec 23 1996 Time Approved : 15:11 Date Received : Oct 09 1996

Time Received: 08:52

GWA charged : 522020702 Reviewed by JACQUIE JANIBAGIAN

MSA mR/hr : None Signature \_\_\_\_\_

Laboratory QA Review Hazard Index : None

Signature \_\_\_\_\_ PCBs >50 ppm : NO

COMMENTS:

	Lab Field	
Analysis	Spl ID Spl ID	Method Analyst Results
Acid	6DC30 WL-104	17012 RAH titrated less than .5 ml
Chloride	6DC30 WL-104	57171 RAH < 1.25947E+01 ug/ml
Fluoride	6DC30 WL-104	87092 RAH 3.49E+01 +- 1.7E+00 ug/ml
SpGr	6DC30 WL-104	47981 BCS 9.99012E-01 +- 2.6E-04 @ 25/4
Sulfate	6DC30 WL-104	97168 BCS < 9.03652E+00 uG/mL
Uranium	6DC30 WL-104	17920 RAH 2.35E-02 +- 1.3E-03 G/L
	6DC30 WL-104	17920 RAH 2.23E-02 +- 1.6E-03 G/L
рН	6DC30 WL-104	87017 RLC 2.75E+00 +- 1.6E-01 pH
End of Report	8 results.	

#### FINAL REPORT for WL-105:105 DEMIN

Log Type: \*\* PLANT \*\*

Log Number : 97-09169 Phone Number : 6-3846 Report for : R L HASTINGS Mailstop : 5111

Date Approved : Sep 29 1997 Time Approved : 17:30 Date Received : Sep 16 1997

Time Received: 22:48

GWA charged : 522020602 Reviewed by ROBERTA JORDAN

MSA mR/hr : None Signature \_\_\_\_

Hazard Index : None Laboratory QA Review

PCBs >50 ppm : NO Signature \_\_\_\_\_

COMMENTS:

Analysis	Lab Field Spl ID Spl ID	Method Analyst Results
Acid	7HR36 SAMPLE	3 87012 SRT <.0241 Normal Acid
Aluminum	7HR36 SAMPLE	3 87100 BCS < 4.74571E-04 MOLAR
	7HR36 SAMPLE	3 87100 BCS < 6.72322E-04 MOLAR
Chloride	7HR36 SAMPLE	3 57171 RAH < 4.11211E+00 ug/mL
Fluoride	7HR36 SAMPLE	3 87092 RAH 5.07E-04 +- 8.1E-05 molar
GROSS BETA	7HR36 SAMPLE	3 87970 KFM 2.05E+03 +- 1.9E+02 B/Min/ml
Sulfate	7HR36 SAMPLE	3 97168 BCS < 4.06935E+00 ug/ml
Uranium	7HR36 SAMPLE	3 17920 KFM < 3.46607E-01 ug/mL
	7HR36 SAMPLE	3 17920 KFM < 3.46607E-01 ug/mL
pН	7HR36 SAMPLE	3 87017 SRT 2.404E+00 +- 5.6E-02

pH 7HR36 SAMPLE 3 End of Report -- 10 results.



#### **COVER PAGE**

1. SDG Transmittal Date: JUN 1 3 2000 2. Subcontractor Name: INTEC ANALYTICAL CHEMISTRY LABORATORY Analytical Laboratories Department Bechtel BWXT Idaho, LLC 3. Contract Number: ER-SOW-169 4. SDG Type: Semivolatile Organics by GC/MS 5. Reporting Tier: Tier II 6. SDG Number: NCD-123 7. SAP Number: N/A ORIGINAL 8. Applicable TOS Modification Numbers: N/A Catherine A. Crowder ALD Organic Analyses Supervisor 12/2000

Patrick D. Troescher

Mariam R. Thomas

ALD Assistant Quality Assurance Officer

Semivolatile Analyses Technical Leader

Date

#### 1D SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET APPENDIX IX ANALYTES

	INEEL Sample No.	NCD-123
Lab Name: Analytical Chemistry Lab	Contract: NA	
Lab Code: ALD-INTEC TOS No: NA	Method No: 9270 SDG N	No: <u>NCD-123</u>
Matrix (soil/water): WATER	Lab Sample ID: 9AH61: 9A	H61DL10
Sample wt/vol: 100 (g/mL) mL	Lab File ID: <u>SS200192</u> ; SS2	200303
Level: (low/med) LOW	Date Received: 02/10/99	····
% Moisture: NA Decanted (Y/N): NA	Date Extracted: 02/11/99	
Concentrated Extract Volume: 1000 (µL)	Date Analyzed: 02/16/99: 0	3/08/99
Injection Volume:1 (µL)	Dilution Factor: 1: 10	
GPC Cleanup (Y/N): NO	pH: <u>NA</u>	

CAS NO.	COMPOUND	CONCENTRATION UNITS 'ug/L	Q
62-75-9	N-Nitrosodimethylamine	250	UD
110-86-1	Pyridine	250	UD
108-95-2	Phenol	250	UD
111-44-4	bis(2-Chloroethyl)ether	250	UD
95-57-8	2-Chlorophenol	250	UD
541-73-1	1,3-Dichlorobenzene	250	UD.
106-46-7	1,4-Dichlorobenzene	250	UD
95-50-1	1,2-Dichlorobenzene	250	UD
95-48-7	2-Methylphenol	250	UD
108-60-1	bis(2-Chloroisopropyl)ether	250	UDM
106-44-5	3 & 4-Methylphenol	250	UD
621-64-7	N-Nitroso-di-n-propylamine	250	UD
67-72-1	Hexachloroethane	250	UD
98-95-3	Nitrobenzene	25	U
78-59-1	Isophorone	25	U
88-75-5	2-Nitrophenol	25	Ŭ
105-67-9	2,4-Dimethylphenol	25	U
111-91-1	bis(2-Chloroethoxy)methane	25	บ
120-83-2	2,4-Dichlorophenol	25	U
120-82-1	1,2,4-Trichlorobenzene	. 25	υ
91-20-3	Naphthalene	25	Ū.
106-47-8	4-Chloroaniline	25	Ū
87-68-3	Hexachlorobutadiene	25	U.

#### 1E SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET APPENDIX IX ANALYTES

INEEL Sample No.	NCD-123
	<u> </u>

••	·
Lab Name: Analytical Chemistry Lab	Contract: NA
Lab Code: ALD-INTEC TOS No: NA	Method No: <u>9270</u> SDG No: <u>NCD-123</u>
Matrix (soil/water): WATER	Lab Sample ID: 9AH61: 9AH61DL10
Sample wt/vol: 100 (g/mL) mL	Lab File ID: <u>SS200192</u> ; <u>SS200303</u>
Level: (low/med) LOW	Date Received: 02/10/99
% Moisture: NA Decanted (Y/N): NA	Date Extracted: 02/11/99
Concentrated Extract Volume: 1000 (µL)	Date Analyzed: 02/16/99 : 03/08/99
Injection Volume:1_(µL)	Dilution Factor: 1; 10
GPC Cleanup (Y/N) NO	nH. NA

CAS NO.	COMPOUND	CONCENTRATION UNITS ug/L	Q
59-50-7	4-Chloro-3-methylphenol	25	U
91-57-6	2-Methylnaphthalene	25	Ü
77-47-4	Hexachlorocyclopentadiene	25	U
88-06-2	2,4,6-Trichlorophenol	25	U
95-95-4	2,4,5-Trichlorophenol	25	U
91-58-7	2-Chloronaphthalene	25	U
88-74-4	2-Nitroaniline	25	U
131-11-3	Dimethylphthalate	25	U
606-20-2	2,6-Dinitrotoluene	25	Ū-
208-96-8	Acenaphthylene	25	υ
99-09-2	3-Nitroaniline	25	U
83-32-9	Acenaphthene	25	U
51-28-5	2,4-Dinitrophenol	76	М
100-02-7	4-Nitrophenol	25	υ
132-64-9	Dibenzofuran	25	Ŭ
121-14-2	2,4-Dinitrotoluene	25	υ
84-66-2	Diethylphthalate	25	ŬМ
7005-72-3	4-Chlorophenyl-phenylether	25	υ
86-73-7	Fluorene	25	Ū
100-01-6	4-Nitroaniline	25	Ŭ
534-52-1	4,6-Dinitro-2-methylphenol	25	Ü
86-30-6	N-Nitrosodiphenylamine	25	Ŭ
126-73-8	Tri-n-butyl phosphate	25	UM

#### 1F SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET APPENDIX IX ANALYTES

	INEEL Sample No.	NCD-123
Lab Name: Analytical Chemistry Lab	Contract: NA	
Lab Code: ALD-INTEC TOS No: NA	Method No: 9270 SDG No: NCD-123	
Matrix (soil/water): WATER	Lab Sample ID: 9AH61: 9AH61DL10	
Sample wt/vol: 100 (g/mL) mL	Lab File ID: <u>SS200192</u> ; <u>SS200303</u>	
Level: (low/med) LOW	Date Received: 02/10/99	
% Moisture: NA Decanted (Y/N): NA	Date Extracted: 02/11/99	
Concentrated Extract Volume: 1000 (µL)	Date Analyzed: 02/16/99; 03/08/99	
Injection Volume:1 (µL)	Dilution Factor: 1: 10	
GPC Cleanup (Y/N): NO	pH: NA	

CAS NO.	COMPOUND	CONCENTRATION UNITS ug/L	Q
103-33-3	Azobenzene	25	U
101-55-3	4-Bromophenyl-phenylether	25	U
118-74-1	Hexachlorobenzene	25	U
87-86-5	Pentachlorophenol	25	Ü
85-01-8	Phenanthrene	25	U
120-12-7	Anthracene	25	U
86-74-8	Carbazole	25	U
84-74-2	Di-n-butylphthalate	25	Ŭ
206-44-0	Fluoranthene	25	Ŭ.
129-00-0	Pyrene	25	U
85-68-7	Butylbenzylphthalate	. 25	U
91-94-1	3,3'-Dichlorobenzidine	25	U
218-01-9	Chrysene	25	U
56-55-3	Benzo(a)anthracene	25	U
117-81-7	bis(2-Ethylhexyl)phthalate	47	
117-84-0	Di-n-octylphthalate	25	U
205-99-2	Benzo(b)fluoranthene	25	ΰ
207-08-9	Benzo(k)fluoranthene	25	U
50-32-8	Benzo(a)pyrene	25	U
193-39-5	Indeno(1,2,3-cd)pyrene	25	U
53-70-3	Dibenzo(a,h)anthracene	25	U
191-24-2	Benzo(g,h,i)perylene	25	U

# SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET TENTATIVELY IDENTIFIED COMPOUNDS APPENDIX IX ANALYTES

	INEEL Sample No.	NCD-123
Lab Name: Analytical Chemistry Lab	Contract: NA	
Lab Code: ALD-INTEC TOS No: NA	Method No: <u>9270</u> SDG No: <u>NCD-123</u>	
Matrix (soil/water): WATER	Lab Sample ID: 9AH61: 9AH61DL10	
Sample wt/vol: 100 (g/mL) mL	Lab File ID: <u>SS200192</u> : <u>SS200302</u>	
Level: (low/med) LOW	Date Received: 02/10/99	
% Moisture: NA Decanted (Y/N): NA	Date Extracted: 02/11/99	
Concentrated Extract Volume: 1000 (µL)	Date Analyzed: 02/16/99; 03/08/99	
Injection Volume:1_(µL)	Dilution Factor: 1: 10	
GPC Cleanup (Y/N): NO pH: N/A	Concentration Units: ug/L	
No. TICs Found: 20		

	CAS NO.	COMPOUND	RT	EST. CONC. ug/L	Q
1		Unknown	8:45	550	ло
2		Unknown	10:49	1900	)D
3		Unknown	10:54	2000	JD
4		Unknown	11:07	2200	JD
5		Unknown	13.97	21	J
6	65-85-0	Benzoic Acid	13.98	21	JN
7		Unknown	14.08	31	J
8		Unknown	15.82	35	J
.9		Unknown	16.52	20	J
10		Unknown	17.50	24	J
11		Unknown	18.22	38	J
12		Unknown	19.27	19	J
13		Unknown	19.87	56	J
14		Unknown	21.43	35	J
15		Unknown	22.94	19	J
16		Unknown	29.32	41	J
17	• • •	Unknown	32.88	320	J
18		Unknown	38.06	820	J
19		Unknown	44.50	1200	J
20		Unknown	52.83	1100	J

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#### **COVER PAGE**

1. SDG Transmittal Date: July 19, 2000	
2. Subcontractor Name: INTEC ANALYTICAL CHEMIS Analytical Laboratories Department Bechtel BWXT Idaho, LLC	
3. Contract Number: ER-SOW-169	, ~
4. SDG Type: Semivolatile Organics by GC/MS	5003013
5. Reporting Tier: Tier II	600,30
6. SDG Number: NCC101/119-000301	
7. SAP Number: N/A	
8. Applicable TOS Modification Numbers: N/A	
Jeffrey L. Jeter	7/19/00 Date
Acting ALD Organic Analyses Supervisor	
Shelly J. Sailer ALD Quality Assurance Officer	07/19/2000 Date
Mariam R. Thomas	7/19/00 Date

ORIGINAL

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Semivolatile Analyses Technical Leader

# 1D SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET APPENDIX IX ANALYTES

INEEL Sample No.

WL-106-000307

Lab Name: Analytical Chemistry Lab	Contract: N/A	
Lab Code: ALD-INTEC TOS No: N/A	Method No: 9270 SDG No: NCC101/119-000301	
Matrix (soil/water): WATER	Lab Sample ID: 0AN16	
Sample wt/vol: 100 (g/mL) mL	Lab File ID: SS201499	
Level: (low/med) LOW	Date Received: 03/08/00	
% Moisture: N/A Decanted (Y/N): N/A	Date Extracted: 03/08/00	
Concentrated Extract Volume:1000_ (µL)	Date Analyzed: 03/20/00	
Injection Volume:1 (µL)	Dilution Factor: 1	
GPC Cleanup (Y/N): NO	pH: <u>N/A</u>	

CAS NO.	COMPOUND	CONCENTRATION UNITS ug/L	Q
62-75-9	N-Nitrosodimethylamine	20	UM
110-86-1	Pyridine	20	U
108-95-2	Phenol	20	U
111-44-4	bis(2-Chloroethyl)ether	20	Ū
95-57-8	2-Chlorophenol	20 .	U
541-73-1	1,3-Dichlorobenzene	20	U
106-46-7	1,4-Dichlorobenzene	20	υ
95-50-1	1,2-Dichlorobenzene	20	U
95-48-7	2-Methylphenol	20	U
108-60-1	bis(2-Chloroisopropyl)ether	20`	U
106-44-5	3 & 4-Methylphenol	20	U
621-64-7	N-Nitroso-di-n-propylamine	20	U
67-72-1	Hexachloroethane	20	U
98-95-3	Nitrobenzene	20	UM '
78-59-1	Isophorone	. 20	U
88-75-5	2-Nitrophenol	20	U
105-67-9	2,4-Dimethylphenol	20	U
111-91-1	bis(2-Chloroethoxy)methane	20	U
120-83-2	2,4-Dichlorophenol	20	U
120-82-1	1,2,4-Trichlorobenzene	20	บ
91-20-3	Naphthalene	20 ·	U
106-47-8	4-Chloroaniline	20	U
87-68-3	Hexachlorobutadiene	20	U

#### 1E SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET APPENDIX IX ANALYTES

INEEL	Sample	No.
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WL-106-000307

Lab Name: _Analytical Chemistry Lab         Lab Code: _ALD-INTEC_ TOS No: _N/A         Matrix (soil/water): _WATER         Sample wt/vol: _100	Contract: N/A  Method No: 9270 SDG No: NCC101/119-000301  Lab Sample ID: 0AN16  Lab File ID: SS201499  Date Received: 03/08/00  Date Extracted: 03/08/00  Date Analyzed: 03/20/00  Dilution Factor: 1
GPC Cleanup (Y/N): NO	pH: <u>N/A</u>

CAS NO.	COMPOUND	CONCENTRATION UNITS ug/L	Q
59-50-7	4-Chloro-3-methylphenol	20	U
91-57-6	2-Methylnaphthalene	20	U
77-47-4	Hexachlorocyclopentadiene	20	U
88-06-2	2,4,6-Trichlorophenol	20	U
95-95-4	2,4,5-Trichlorophenol	20	U
91-58-7	2-Chloronaphthalene	. 20.	U
88-74-4	2-Nitroaniline	20	UM
131-11-3	Dimethylphthalate	20	U
606-20-2	2,6-Dinitrotoluene	20	U
208-96-8	Acenaphthylene	20	Ŭ
99-09-2	3-Nitroaniline	20	U
83-32-9	Acenaphthene	20	U
51-28-5	2,4-Dinitrophenol	13	J
100-02-7	4-Nitrophenol	20	U
132-64-9	Dibenzofuran	20	U
121-14-2	2.4-Dinitrotoluene	20	U
84-66-2	Diethylphthalate	20	UM
7005-72-3	4-Chlorophenyl-phenylether	20	U
86-73-7	Fluorene	20	U
100-01-6	4-Nitroaniline	20	U
534-52-1	4,6-Dinitro-2-methylphenol	5	J
86-30-6	N-Nitrosodiphenylamine	20	U
126-73-8	Tri-n-butyl phosphate	170	EM

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### 1F SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET APPENDIX IX ANALYTES

INEEL Sample No.

WL-106-000307

Lab Name: Analytical Chemistry Lab  Lab Code: ALD-INTEC TOS No: N/A  Matrix (soil/water): WATER  Sample wt/vol: 100 (g/mL) mL  Level: (low/med) LOW  % Moisture: N/A Decanted (Y/N): N/A  Concentrated Extract Volume: 1000 (µL)	Contract: N/A  Method No: 9270 SDG No: NCC101/119-000301  Lab Sample ID: 0AN16  Lab File ID: SS201499  Date Received: 03/08/00  Date Extracted: 03/08/00  Date Analyzed: 03/20/00  Dilution Factor: 1
Injection Volume: 1 (μL)  GPC Cleanup (Y/N): NO	Dilution Factor: _1pH: _N/A

ip (Y/N): <u>NC</u> ————————————————————————————————————	COMPOUND	· CONCENTRATION UNITS ug/L	Q
CAS NO.		20	UM
103-33-3	Azobenzene	20	, U
101-55-3	4-Bromophenyl-phenylether	20	U
118-74-1	Hexachlorobenzene	20	U
87-86-5	Pentachlorophenol	20	U
85-01-8	Phenanthrene	20	U
120-12-7	Anthracene	20	U
86-74-8	Carbazole	20	UM
84-74-2	Di-n-butylphthalate	20	U
206-44-0	Fluoranthene	20	U
129-00-0	Pyrene	20	U
85-68-7	Butylbenzylphthalate	20	U
91-94-1	3,3'-Dichlorobenzidine	20	U
218-01-9	Chrysene		U
56-55-3	Benzo(a)anthracene	20	<del>                                     </del>
117-81-7	bis(2-Ethylhexyl)phthalate	31	U
117-84-0	Di-n-octylphthalate	20	U
205-99-2	Benzo(b)fluoranthene	20	T <sub>U</sub>
207-08-9	Benzo(k)fluoranthene	20	T U
50-32-8	Benzo(a)pyrene	. 20	U
193-39-5	Indeno(1,2,3-cd)pyrene	20	<u>U</u>
53-70-3	Dibenzo(a,h)anthracene	20	<del>  U</del>
191-24-2	Benzo(g,h,i)perylene	20	

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#### 1M SEMIVOLATILE ORGANICS ANALYSIS DATA SHEET TENTATIVELY IDENTIFIED COMPOUNDS APPENDIX IX ANALYTES

INEEL	Sample	No.
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WL-106-000307

ab Code: ALD-INTEC TOS No: N/A  Matrix (soil/water): WATER	Contract: N/A  Method No: 9270 SDG No: NCC101/119-00030 1  Lab Sample ID: 0AN16  Lab File ID: SS201499  Date Received: 03/08/00
_evel: (low/med) _LOW	Date Extracted: 03/08/00  Date Analyzed: 03/20/00  Dilution Factor: 1
Injection Volume:	pH: <u>NA</u>

Г	CAS NO.	COMPOUND	RT (min)	EST. CONC. ug/L	Q
<del>.  -</del>		Unknown	3:56	14	J
1		Unknown	5:57	11	J
2	: (7. (6.2	Chloroform	4:17	12	J
3	67-66-3	Unknown	7:28	85	J
4		Unknown	13:06	6300	J
5		Unknown	16:10	350	J
6		Substituted benzene	21:36	37	J
7		Substituted benzene	21:56	23	J
8		Substituted benzene	22:00	24	J
9		Substituted benzene	22:12	23	J
10		Substituted benzene	22:32	28	J
11		Substituted benzene	23:20	24	1
12		Substituted benzene	23:26	14	J
13		Substituted benzene	23:38	13	J
14		Substituted benzene	23:59	11	J
15			24:07	19	ЛВ
16		Unknown	24:30	27	J
17		Substituted benzene	28:44	21	J
18		Unknown	29:31	22	1
19		Unknown	30:18	65	1
20		Unknown	30.16		

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